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FILE COVERS 1961 - 9 Jul 2002 VOL 1-7 ISS 1

FILE LAST UPDATED: 9 Jul 2002 (20020709/ED)

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word all tot 179

139 ANSWER 1 OF 14 HCAPLUS COPYRIGHT 2002 ACS

AN 1003:491818 HCAPLUS

EN 136:382505

TI Device for monitoring cells

IN Eitner, J. Bruce; Hemperly, John Jacob; Guarino, Richard D.; Wodnicka, Magdalena; Stitt, David T.; Burrell, Gregory J.; Foley, Timothy G., Jr.; Beatty, Patrick Shawn

PA **Becton, Dickinson and Company, USA**

SO U.S., 41 pp., Cont.-in-part of U.S. Ser. No. 715,127.

CODEN: USXXAM

DT Patent

LA English

IC ICM C13Q901-11

NCL 435032000

CC 9-1 (Biochemical Methods)

Section cross-references : 1, 4

FAN.CHT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 6196506	B1	1999-08-23	US 1999-042720	1999-06-29
	EP 509791	A1	1999-10-01	EP 1999-305891	1999-04-15
	EP 509791	B1	1999-07-03		
	E: DE, FR, GB, IT				
	CA 2060329	AA	1999-10-19	CA 1999-0160329	1999-10-16
	JP 35137596	A1	1999-06-01	JP 1999-01308	1999-10-13
	JP 07073510	B4	1999-08-19		
PRAI	US 1991-087399	B1	1991-04-15		
	US 1991-25899	A1	1991-03-15		
	US 1990-11557	A1	1990-10-15		

AB The present invention relates to methods for detection and evaluation of metabolic activity of eukaryotic and/or prokaryotic cells based upon their ability to consume dissolved oxygen. The methods utilize a **luminescence** detection system which makes use of the sensitivity

of the **luminescent** emission of certain compds. to the presence of oxygen, which quenches (diminishes) the compd.'s **luminescent** emission in a **concn.** dependent manner. Respiring eukaryotic and/or prokaryotic cells will affect the oxygen **concn.** of a liq. medium in which they are immersed. Thus, this invention provides a convenient system to gather information on the presence, identification, quantification and cytotoxic activity of eukaryotic and/or prokaryotic cells by detg. their effect on the oxygen **concn.** of the media in which they are present.

ST Device monitoring cell

IT Plates

(Microtitration; device for monitoring cells)

IT **Analytical apparatus**

Antibiotics

Bacteria (Eubacteria)

Biological materials

Blood

Blood serum

Cell

Cell proliferation

Chemicals

Coating materials

Composition

Concentration (condition)

Culture media

Cytotoxicity

Drugs

Escherichia coli

Eukaryote

Extracellular matrix

Fluorescence quenching

Impermeability

Insecta

Light

Liquids

Luminescence

Luminescence quenching

Luminescence spectroscopy

Luminescent substances

Mathematical methods

Metabolism

Microorganism

Molecules

Particles

Permeability

Prokaryote

Pseudomonas aeruginosa

Radiation

Reducing agents

Respiration, animal

Respiration, microbial

Sensors

Solutes

Wavelength

Wetting

Yeast

(device for monitoring cells)

IT Toxins

PL: ADV (Adverse effect, including toxicity); BIOL (Biological study)

(device for monitoring cells)

IT Reagents

PL: ARS (Analytical reagent use); ANST (Analytical study); USES (Uses)

(device for monitoring cells)

IT Plastics, analysis
FL: AFU (Analytical role, unclassified); ANST (Analytical study)
(device for monitoring cells)

IT Rubber, analysis
FL: AFU (Analytical role, unclassified); ANST (Analytical study)
(device for monitoring cells)

IT Silicone rubber, analysis
FL: AFU (Analytical role, unclassified); ANST (Analytical study)
(device for monitoring cells)

IT **Growth factors, animal**
FL: BSU (Biological study, unclassified); BIOL (Biological study)
(device for monitoring cells)

IT Collagens, biological studies
FL: BSU (Biological use, unclassified); BIOL (Biological study ; USES
(Uses)
(device for monitoring cells)

IT Entactin
FL: BSU (Biological use, unclassified); BIOL (Biological study); USES
(Uses)
(device for monitoring cells)

IT Laminins
FL: BSU (Biological use, unclassified); BIOL (Biological study); USES
(Uses)
(device for monitoring cells)

IT Proteoglycans, biological studies
FL: BSU (Biological use, unclassified); BIOL (Biological study); USES
(Uses)
(heparitin sulfate-contg.; device for monitoring cells)

IT **Optical detectors**
(luminescence; device for monitoring cells)

IT Animal cell
(mammal; device for monitoring cells)

IT Amino acids, biological studies
FL: BSU (Biological use, unclassified); BIOL (Biological study); USES
(Uses)
(nonessential; device for monitoring cells)

IT Collagens, biological studies
FL: BSU (Biological use, unclassified); BIOL (Biological study); USES
(Uses)
(type IV; device for monitoring cells)

IT 1494-18-1, 9,10-Diphenylanthracene 18153-62-0D, Tris-2,2'-
bipyridylruthenium (II), salts 36309-38-3, Tris-4,7-diphenyl-1,10-
phenanthroline ruthenium (II) chloride 59525-27-4, Tris-2,2'-
bipyridylruthenium (II) chloride hexahydrate. 63373-04-6D,
Tris-4,7-diphenyl-1,10-phenanthroline ruthenium (II), salts
FL: ARU (Analytical reagent use); ANST (Analytical study); USES (Uses)
(device for monitoring cells)

IT 7631-86-3, Silica, analysis
FL: AFU (Analytical role, unclassified); ANST (Analytical study)
(device for monitoring cells)

IT 59-05-1, Methotrexate 151-21-3, Sodium dodecyl sulfate, biological
studies 365-21-4, Vinblastine 7757-33-3, Sodium Sulfite 7782-44-7,
Oxygen, biological studies 26623-22-8, Sodium Azide 35607-56-0,
Cefixitin 89263-75-2, Cefuroxime 55721-33-1, Ciprofloxacin
FL: BSU (Biological study, unclassified); BIOL (Biological study)
(device for monitoring cells)

IT 57-12-1, Streptomycin, biological studies 113-24-6, Sodium pyruvate
1397-34-3, Fungizone 1406-95-3, Penicillin 119978-13-6, Matrigel
141907-41-7, Matrix metalloproteinase
FL: BSU (Biological use, unclassified); BIOL (Biological study ; USES
(Uses)
(device for monitoring cells)

RE.CNT 1 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD

PE

- (1) Bacon, J; Anal Chem 1987, V59(23), P2759 HCAPLUS
- (2) Berndt; US 6395574 A 1990
- (3) Collins; US 6107082 A 1990
- (4) Gentile; US 5938517 A 1999 HCAPLUS
- (5) Goswami, K; Fiber Optic Chemical Sensor for the Measurement of Partial Pressure of Oxygen 1988, V890, P111
- (6) Stitt; US 5567398 A 1996
- (7) Walt; US 5244636 A 1993 HCAPLUS
- (8) Wertz; US 4448134 A 1984
- (9) Wilfobeis, O; Mikrochimica Acta 1986, V3(5-6), P835 HCAPLUS

L79 ANSWER 1 OF 14 HCAPLUS COPYRIGHT 2002 ACS

AN 2002:164471 HCAPLUS

DI 136:196648

TI Method and apparatus for non-destructive screening of clinical specimen integrity

IN Samsundar, James; Jacobs, Merritt Nyles

PA One Telometrix Inc., San.

SO U.S., 21 pp., Cont.-in-part of U. S. Ser. No. 541,390, abandoned.

CODEN: USXXAM

DT Patent

LA English

IC ICM G01N033-48

NCL B56040000

CI 9-1 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 6395471	B1	20020305	US 1997-081606	19970609 <--
PRAI	US 1996-041340	B2	19961010	<--	

AB A method and app. for providing a non-destructive pre-test screen of specimen integrity for a blood analyzer by measurement of absorbance or reflectance is provided. The method involves measurement of polychromatic **light** in the near IR and adjacent visible region, which is either transmitted or reflected from a specimen as presented for measurement, and correlation of the measurement, on the basis of predetd. **algorithms**, to the quantity of a known substance contained in the sample. The app. employs a spectrophotometer which emits radiation which is split into a beam which passes to a sample and a ref. beam, the beam returning from the sample and the ref. beam are variably combined and further sepd. into various components by means of a grating and focused onto a linear array detector. A microprocessor receives output from the array detector and performs **calcns.** of **concn.** (s) of the known substance(s). The invention provides quality assurance for state-of-the-art blood analyzers and **automated** labs. by pre-screening serum and plasma integrity, even where labels on the sample container would normally interfere with a quality assurance assessment, identifying samples not suitable for certain blood tests, or, if tests are conducted on specimens with compromised integrity, the pre-screening results will aid in the interpretation of the test results.

ST app screening clin specimen.

IT **Sensors**

(linear array; method and app. for non-destructive screening of clin. specimen integrity)

IT **Light**

(Polychromatic; method and app. for non-destructive screening of clin. specimen integrity)

IT **Analysis**(clin.; method and **app.** for non-destructive screening of clin. specimen integrity)

IT Absorption spectroscopy

Algorithm

Analytical apparatus

Blood analysis

Blood plasma

Blood serum

Concentration (condition)

Containers

Diffraction gratings

Frequency

Labels

Light**Mathematical methods**

Molecules

Optical reflection**Optical transmission**

Quality control

Radiation

Samples

Spectra

Spectrometers

Standard substances, analytical

Time**Turbidity**

UV and visible spectroscopy

Wavelength

(method and app. for non-destructive screening of clin. specimen integrity)

IT Hemoglobins

EL: ANT (Analyte); ANST (Analytical study)

(method and app. for non-destructive screening of clin. specimen integrity)

IT **Computers**

(microprocessors; method and app. for non-destructive screening of clin. specimen integrity)

IT **IR radiation**

IR spectroscopy

(near-IR; method and app. for non-destructive screening of clin. specimen integrity)

IT Soybean oil

EL: AMI (Analyte); ANST (Analytical study)

(phospholipid-stabilized; method and app. for non-destructive screening of clin. specimen integrity)

IT 114-25-0, Biliverdin 635-65-4, Bilirubin, analysis

EL: ANT (Analyte); ANST (Analytical study)

(method and app. for non-destructive screening of clin. specimen integrity)

RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD

FE

(1) Anon; CA 2019011 1994

(2) Coggeshall; US 3701736 A 1972

(3) Heinemann; US 5291884 A 1994

(4) Jacobs; US 5846492 A 1998

(5) Jacques; US 5313791 A 1994

(6) Karkar; US 5066459 A 1991

(7) Lindsaard; US 5281646 A 1994

(8) Lindsaard; US 5866903 A 1994

(9) McNeal; US 5734468 A 1998

(10) Petratz; US 5351685 A 1994

(11) Parry; US 5360904 A 1994

L'9 ANSWER 3 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 2001:585529 HCAPLUS

DN 136:17937

TI System and method for analyzing antibiotic susceptibility of biological

samples
 IN Wiles, Timothy M.; Turner, David J.; O'Connell,
 Michael A.; Parmigiani, Giovanni; Clyde, Merlise
 PA Becton, Dickinson and Co., USA
 SO Eur. Pat. Appl., 32 pp.
 CODEN: EPEXDEW
 DT Patent
 LA English
 IC ICM G01N021-31
 CC 10-5 (Microbial, Algal, and Fungal Biochemistry)
 Section cross-reference(s): 1

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 1160564	A2	200111295	EP 2001-111411	20010510 <--
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, EO				
	JP 2001121697	A2	20020519	JP 2001-160395	20010529 <--
PRAI	US 2000-081891	A	20000131	<--	
AB	A system and method for analyzing samples, such as biol. samples, to accurately and effectively det. the susceptibility of the samples to antimicrobial materials, to det. min. inhibitory concn. (MIC) values for the resp. samples and antimicrobial materials. At each of a plurality of time intervals, the system and method directs a plurality of different analyzing light wavelengths, such as red, green and blue wavelengths, onto each of a plurality of sample wells, and detects a resp. resultant light wavelength emanating from the resp. sample wells for each of the analyzing light wavelengths. The system and method uses resultant light wavelengths to generate at least two growth indicator characteristic curves representing, for example, the redox state and turbidity characteristics of the sample wells. The system then uses the redox state and turbidity characteristics of sample wells contg. the same antimicrobial material to det. the MIC value for that material in relation to the sample contained in those wells.				
ST	antibiotic susceptibility biol sample				
IT	Antibiotics Antimicrobial agents Computer application Drug screening Mathematical methods Measuring apparatus Redox potential Turbidity (system and method for analyzing antibiotic susceptibility of biol. samples)				

L19 ANSWER 4 OF 14 HCAPLUS COPYRIGHT 2002 ACS

AN 2001:8170:8 HCAPLUS

DN 135:128025

TI Method for non-invasive spectrophotometric blood oxygenation monitoring

IN Berni, Paul

PA Gas Medical Systems, Inc., USA

SO PCT Int. Appl., 36 pp.

CODEN: PEXKDE

DT Patent

LA English

IC ICM G01N

CC 9-1 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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Oxygen; method for non-invasive spectrophotometric blood oxygenation monitoring;

IT 7781-44-7, Oxygen, analysis

PL: ANT (Analyte); ANST (Analytical study)

sensors; method for non-invasive spectrophotometric blood oxygenation monitoring;

L79 ANSWER 1 OF 24 HCAPLUS COPYRIGHT 2001 ACS

AN 2001:040972 HCAPLUS

DN 123:177039

TI Method and apparatus for determining the sensitivity of a microorganism to a **growth** altering agent

IN Wardlaw, Stephen C.

PA USA

SO U.S., 4 pp., Cont.-in-part of U.S. 6,022,734.

CODEN: USKXAM

DT Patent

LA English

IC ICM 01LM001-16

ICS 01SQ001-18

NCL 415286700

CC 9-1 (Biochemical Methods)

Section cross-reference(s): 1, 10

FAN.CNT 7

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FI	US 6234516	B1	20010904	US 1116-477932	10000105 <--
	US 6011714	A	20000308	US 1999-256451	19990223 <--
	US 6140069	A	20001031	US 1999-256651	19990223 <--
	NO 2000084456	A	20001008	NO 1999-4459	10000306 <--
	NO 2001030043	A	20010706	NO 1991-45	10010104 <--
	EP 1120467	A	20010601	EP 2011-300967	10010105 <--
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
	JP 2001114597	A1	20010814	JP 2001-330	10010105 <--
	CN 1173986	A	20011108	CN 2011-117354	10010105 <--
PFAI	US 1998-772169	P	19980307	<--	
	US 1998-772171	P	19980307	<--	
	US 1999-256651	A1	19990223	<--	
	US 1999-256451	A1	19990223	<--	
	WO 1999-274511	W	19990308	<--	
	US 2000-477932	A	20000105	<--	

AB A method and an app. for detg. the **concn.** at which a **growth**-altering agent has an appreciable effect on the **growth** of a target microorganism are provided. The method comprises the steps of (a) providing a microorganism **growth** medium; (b) providing a sensible reagent, which includes a **growth**-altering agent mixed with a marker that has a signal with a magnitude proportional to the **concn.** of the marker; (c) incorporating the sensible reagent into the **growth** medium, in a manner that creates a gradient of **growth**-altering agent and marker **concns.** within the **growth** medium; (d) inoculating the **growth** medium with the target microorganism; (e) incubating the inoculated **growth** medium for a period of **time** sufficient for the target microorganism to **grow** a detectable amt.; (f) evaluating **growth** characteristics of the microorganism in a region contg. the **growth**-altering agent; (g) measuring the magnitude of the marker signal in that region; and (h) detg. the **concn.** of the **growth**-altering agent using the measured magnitude of the marker signal.

ST app detg microorganism **growth** agent

IT Molecules

(**Growth** altering; method and app. for detg. sensitivity of a

microorganism to a **growth** altering agent)

IT **Fluorometers**

(Scanning; method and app. for detg. sensitivity of a microorganism to a **growth** altering agent)

IT Antimicrobial agents

Apparatus

Concentration (condition)

Culture media

Growth, microbial

Light scattering

Mathematical methods

Microorganism

Mixing

Sensors

Time

(method and app. for detg. sensitivity of a microorganism to a **growth** altering agent)

IT Reagents

AL: AFG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(method and app. for detg. sensitivity of a microorganism to a **growth** altering agent)

RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) Anon; EP 0635120 B1 1999 HCAPLUS
- (2) Baer; US 5547849 1996 HCAPLUS
- (3) Brume; US 5925166 1975
- (4) Ericsson; US 4774758 1988
- (5) Ericsson; US 4625522 1991 HCAPLUS
- (6) Ericsson; US 5634612 1997
- (7) Kjellander; US 4234045 1980
- (8) Lancaster; US 5501952 1996 HCAPLUS
- (9) McCoy; US 5700084 1997 HCAPLUS
- (10) Nasen; US 4730647 1988
- (11) Nishimura; US 5417059 1995
- (12) Robertson; US 5206151 1993 HCAPLUS
- (13) Schalkowsky; US 4514495 1985 HCAPLUS
- (14) Schalkowsky; US 5246837 1993
- (15) Schalkowsky; US 5563843 1996 HCAPLUS
- (16) Smith; US 4950455 1990
- (17) Thompson; US 5164301 1992 HCAPLUS
- (18) Vesterberg; US 4034430 1977 HCAPLUS
- (19) Wardlaw; US 6012784 2000 HCAPLUS
- (20) Wardlaw; US 6140069 2000 HCAPLUS

INT ANSWER 3 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1991:048649 HCAPLUS

IN 155:177713

TI Method for extending the range of an immunoassay

IN Wei, Tie Guan; Panaratz, Thomas John; Chu, Victor Pichai

PA Becton Dickinson Inc., USA

SO U.S., 13 pp., Cont.-in-part of U.S. Ser. No. 166,026, abandoned.

CODEN: USXXAM

DT Patent

LA English

IC ICM G01N033-53

ICS G01N033-543; G01N021-00; A61K049-00; C07K016-01

NCL 435G07100

CC 4-10 (Biochemical Methods)

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 6284472	B1	20010904	US 1999-294489	19990420 <--

FEAI US 1998-156026 B2 19981005 <--

A5 Calibrating an immunoassay by generating two reaction rate measuring curves, from samples having higher and lower relative levels of antigen, extrapolating a combination of the curves to cover sample **concns** . known to contain an excess of antigen relative to an amt. of capture reagent and combining the low end linear portion of the higher reaction rate measuring curve with the higher end portion of the extrapolated reaction rate measuring curve, thereby eliminating measuring inaccuracies otherwise arising from the hook effect. For antigen **concns**. higher than the assay range, a high antigen signal utilizing the two rates avoids reporting false results.

ST extending range immunoassay

IT Proteins, specific or class

EL: ANT (Analyte); ANST (Analytical study)

(C-reactive; method for extending range of immunoassay)

IT Reagents

EL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(Capture; method for extending range of immunoassay)

IT Calibration

Concentration (condition)

Immunoassay

Mathematical methods

Reaction kinetics

Regression analysis

Samples

Volume

(method for extending range of immunoassay)

IT Antigens

EL: ANT (Analyte); ANST (Analytical study)

(method for extending range of immunoassay)

IT Immunoassay

(**turbidimetric**; method for extending range of immunoassay)

REL.CNT 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD

FE

- (1) Chadney; US 5514529 1996 HCAPLUS
- (2) Craple; US 4549661 1986 HCAPLUS
- (3) Diamandis; US 5087413 1992 HCAPLUS
- (4) Frengen; US 5211346 1996 HCAPLUS
- (5) Frengen; US 5733641 1998 HCAPLUS
- (6) Graham; US 4745841 1988 HCAPLUS
- (7) Hirai; US 5348668 1994 HCAPLUS
- (8) Kappe; US 4655752 1977 HCAPLUS
- (9) Kasper; US 4966822 1990 HCAPLUS
- (10) Lindner; US 5578241 1996
- (11) Oh; US 5583055 1996 HCAPLUS
- (12) Oh; US 5703357 1998 HCAPLUS
- (13) Rodriguez; US 4119113 1979 HCAPLUS
- (14) Scatelli; US 5421300 1995 HCAPLUS
- (15) Schaefer; US 5420041 1995 HCAPLUS
- (16) Tung; US 4788133 1988 HCAPLUS
- (17) Wu; US 4858802 1989
- (18) Yamada; US 5353336 1993

L79 ANSWER 7 OF 11 HCAPLUS COPYRIGHT 2002 ACS

AN 2001:095473 HCAPLUS

DI 135:149576

T1 **Automated** optical reader for **multiple** samples, especially for nucleic acid assays

IN Andrews, Jeffrey P.; O'Keefe, Christian V.; Scrivens, Brian G.; Pope, Willard C.; Hansen, Timothy; Failing, Frank

PA **Becton, Dickinson and Company, USA**

SO Eur. Pat. Appl., 46 pp.

CODEN: EPKXDW

DT Patent
 LA English
 IC ICM G01N021-64
 CC 4-1 (Biochemical Methods)
 Section cross-reference(s): 3

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 1134126	A2	20010816	EP 2000-113062	20001221
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, ET, RO				
	JP 2001265272	A2	20010911	JP 2001-6091	20010115
PRAI	US 2000-481686	A	20000114		
AB	<p>An app. and method employ a plurality of light emitting devices which each can get light through a resp. optical fiber toward a resp. sample of a plurality of samples in a time-staggered manner. Light is generated in each of the samples at different times consistent with the times at which light is irradiated onto the sample. A single detector is used to detect the lights emitted from the plurality of samples at these different times. A plurality of bifurcated optical cable are coupled to the light emitting devices and single light detector, and the integrated end of each bifurcated cable acts as the light emitting port and light detecting port. Multiple targets can be detected from each of the plurality of samples in the same manner by providing an app. and method employing a different plurality of light emitting devices and single detector for each target to be detected.</p>				
ST	automated optical reader app nucleic acid assay;				
	multiple sample automated analysis app				
IT	Samples				
	(anal. of multiple ; automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Analysis				
	Process automation				
	(automated anal.; automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Algorithm				
	Electroluminescent devices				
	Fluids				
	Light sources				
	Optical cables				
	Optical detectors				
	Optical fibers				
	Photomultipliers				
	(automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Nucleic acids				
	EL: ANT (Analyte); ANST (Analytical study)				
	(automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Probes (nucleic acid)				
	EL: AFG (Analytical reagent use); ANST (Analytical study); USES (Uses)				
	(automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Analytical apparatus				
	(automated; automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Computers				
	(microcomputers; automated optical reader for multiple samples, esp. for nucleic acid assays)				
IT	Microtiter plates				

(microwell arrays; automated optical reader for
multiple samples, esp. for nucleic acid assays)

L79 ANSWER 8 OF 14 HCAPLUS COPYRIGHT 2002 ACS
AN 2001:417242 HCAPLUS
DN 135:16356
TI Method of measuring tissue hemoglobin saturation using gaussian
decomposition
IN Wilson, David A.
PA Johns Hopkins University, USA
SO PCT Int. Appl., 76 pp.
CODEN: PEXKDL
DI Patent
LA English
IC ICM G01N021-35
ICS A61B005-06
CC 4-5 (Biochemical Methods)
FAN.CMT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FI	WO 2001040770	A1	20010607	WO 2000-0510830	20001204 <--
	W:		AE, AG, AL, AM, AT, AU, AV, BA, BB, BC, BE, BY, BZ, CA, CH, CN,		
			CR, CU, CE, DE, DK, DM, DS, EE, ES, FI, GE, GU, HE, GR, GM, HR,		
			HU, IB, IL, IN, IS, JP, KE, KG, KP, KR, KU, LA, LK, LR, LS, LT,		
			LU, LV, MA, MD, MG, MK, MN, MW, ME, MO, NC, NL, PL, PT, RO, RU,		
			SD, SE, SG, SI, SK, SM, TL, TM, TR, TT, TE, UA, UG, US, UZ, VN,		
			YU, ZA, ZW, AM, AZ, BY, EG, ES, MD, RU, TJ, TM		
	FW:		GH, GM, KE, LS, MW, MC, SD, SL, SS, TE, UG, ZW, AT, BE, CH, CY,		
			DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,		
			BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, ME, SG, TD, TG		

PRAI US 1009-16521P P 19991002 <--

AB The constituents of cerebral tissues that contribute to **light** absorbency, i.e., oxyHb, deoxyHb, water, lipid, cytochrome oxidase and a component for characterizing **light** loss due to scattering, are further characterized and used to construct a model system that emulates cerebral tissue reflectance spectra in a variety of conditions. Using this model system in a reverse mode, compd. spectra collected from brain tissue are de-compd. into individual spectra features. The values for features attributable to oxyHb and deoxyHb are then used to construct a ratio that quantifies the percentage of total Hb that contains oxygen. Because the major portion of **light**, collected by the detecting element of the equipment has transited through brain tissue, this ratio becomes a quant. measure of brain tissue Hb satn. The decompn. anal. method is generally applicable to a variety of tissues besides brain tissue.

ST tissue Hb satn gaussian decompn

IT Energy

(**Light**; method of measuring tissue Hb satn. using gaussian decompn.)

IT Animal tissue

Apparatus

Brain

Databases

Light

Light scattering

Mathematical methods

optical absorption

oxygenation

Reflection spectra

(method of measuring tissue Hb satn. using gaussian decompn.)

IT Hemoglobins

Hemoglobins, oxyhemoglobins

EL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical

study); BIOL (Biological study)
 (method of measuring tissue Hb satn. using gaussian decompn.)
 IT Lipids, biological studies
 PL: KSU (Biological study, unclassified); PEP (Properties); BIOL
 (Biological study)
 (method of measuring tissue Hb satn. using gaussian decompn.)
 IT IR spectroscopy
 (near-IR; method of measuring tissue Hb satn. using gaussian decompn.)
 IT 7782-44-7, Oxygen, analysis
 PL: ANT (Analyte); KSU (Biological study, unclassified); PEP (Physical,
 engineering or chemical process); ANST (Analytical study); BIOL
 (Biological study); PROC (Process)
 (method of measuring tissue Hb satn. using gaussian decompn.)
 IT 4712-18-5, Water, biological studies 9001-16-1, cytochrome oxidase
 PL: KSU (Biological study, unclassified); PEP (Properties); BIOL
 (Biological study)
 (method of measuring tissue Hb satn. using gaussian decompn.)
 PE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
 FE

- (1) Berni, P; PROCEEDINGS OF THE ANNUAL NORTHEAST BIOENGINEERING CONFERENCE
 1998, VCONF 11, P165
- (2) Gieve, E; TEXTILVEREDLUNG 1999, V30(7/93), P169
- (3) Doppel, J; US 5743162 A 1999
- (4) Mannheimer, P; US 5742247 A 1999
- (5) Mather, S; PHYSICS IN MEDICINE AND BIOLOGY 1994, V39(3), P1295
- (6) Schermer, R; APPLIED SPECTROSCOPY 1999, V55(1), P22 HOAPLUS

L79 ANSWER 3 OF 24 HOAPLUS COPYRIGHT 2012 ACS

AN 2001:376867 HOAPLUS

EN 134:55035

TI Method and scattered **light**-measuring apparatus for measuring a
 scattered **light** and method of urinalysis using the same

IN Kawamura, Tatsuaki

PA Matsushita Electric Industrial Co., Ltd., Japan

SO Eur. Pat. Appl., 11 pp.

COVEN: EPXNDW

DT Patent

LA English

IC ICM G01N021-51

CC G-1 (Biochemical Methods)

Section cross-reference(s): 73

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 1102059	A1	20010813	EP 2000-125003	20001118
	E: AU, BR, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
	JP 2001-05046	A1	20010803	JP 2000-321074	20001020 <--
PAI	JP 1999-312780	A	19991113	<--	

AB The present invention provides a method and an app. which eliminate the
 influence of a scattered **light** arising due to the pollutants
 inside and on the surfaces of an optical window, differences in refractive
 index and **light** transmittance of a soln. to be detected, and the
 obstruction due to suspending particles and the like to achieve a
 measurement with high precision and high practicability in the measurement
 of the scattered **light**. The scattered **light**
 propagating within a prescribed angle perpendicularly to the direction of
 propagation of the **light** to be propagated through the inside of
 the soln. is measured. Further, the position of the optical axis of the
light to be propagated through the inside of the soln. and/or the
 position of the photosensor in the direction of the optical axis are set
 so that the influence of the scattered **light** arising at and on
 the surface of the optical axis is not more than a predetd. value within a

practically allowable range. The protein **concn.** in urine was detd. by measuring **turbidity** after heat treatment.

- ST scattered **light** measuring app. urinalysis; protein urine scattered **light** analysis
- IT Proteins, general, analysis
 FL: ANT (Analyte); PEP (Physical, engineering or chemical process); ANST (Analytical study); PROC (Process)
 (coagulation in urine for detg. protein **concn.**; method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)
- IT Particles
 (interfering; method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)
- IT UV and visible spectroscopy
 (light-scattering; method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)
- IT **Light scattering**
 Mathematical methods
 Polarized light
 Refractive index
 Urine analysis
 (method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)
- IT Reagents
 FL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
 (method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)
- IT **Optical instruments**
 (scatterometers; method and scattered **light**-measuring app. for measuring scattered **light** and method of urinalysis using same)

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) Barber, D; US 5040178 A 1991 HCAPLUS
- (2) Canon KK; EP 0459125 A 1991 HCAPLUS
- (3) de Maeyer Leo, G; US 4070410 A 1978
- (4) Kiwa Co; EP 0361771 A 1990

L79 ANSWER 18 OF 14 HCAPLUS COPYRIGHT 2002 ACS

AI: 2000:807747 HCAPLUS

DI: 199:331765

TI: Device and procedure for the monitoring and control of microorganism populations in biologically active fluids

II: Roefler, Thomas; Holzhauser, Peter; Walitza, Eckehard

PA: Fraunhofer-Gesellschaft zur Förderung der Angewandten Forschung EV, Germany

SO: Ger. Offen., 12 pp.

CODEN: GWKXBX

DT: Patent

LA: German

IC: ICM 0100001-02

CC: 9-1 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE 10031999	A1	20001116	DE 1999-19921999	19990512 ---
	WO 2000070078	A2	20001123	WO 2000-EP4289	20000512 ---
	WO 2000070078	A3	20010301		
	W: CA, JP, US				
	RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				

EP 1179174 A2 20020214 EP 2000-936736 20000512 <--
 P: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
 IE, FI

PRAI DE 1999-19921999 A 19990911 <--
 WO 2000-EP4289 W 20000911 <--

AB A data acquisition mechanism and subsequent data processing are used to monitor metabolic parameters in a biol. active fluid so as to detrn. the concn. of organisms. Thus, metabolic products or substrates such as carbon dioxide, hydrogen, oxygen, etc., may be monitored to define bacterial or fungal populations in fluids.

ST microorganism monitoring metabolite liq control app

IT Algorithm

Bacillus (bacterium genus)

Bacillus subtilis

Candida

Clostridium

Control apparatus

Data processing

Desulfotomaculum

Electric conductivity

Enterobacteriaceae

Enterococcus

Escherichia coli

Lactobacillus

Leuconostoc

Liquids

Methanobacterium

Methanococcus

Micrococcus

Pseudomonas

Pseudomonas aeruginosa

Redox potential

Saccharomyces

Sarcina

Sensors

Staphylococcus

Streptococcus

pH

(device and procedure for monitoring and control of microorganism populations in biol. active fluids)

IT Computers

(microprocessors; device and procedure for monitoring and control of microorganism populations in biol. active fluids)

IT Aerobic bacteria

(spore-forming; device and procedure for monitoring and control of microorganism populations in biol. active fluids)

IT 60-21-6, Lactic acid, analysis 64-17-5, Ethanol, analysis 64-18-6, Formic acid, analysis 64-19-7, Acetic acid, analysis 67-64-1, Acetone, analysis 71-23-8, Propanol, analysis 71-36-5, Butanol, analysis 74-82-8, Methane, analysis 107-92-6, Butyric acid, analysis 124-38-9, Carbon dioxide, analysis 133-74-0, Hydrogen, analysis 3312-32-6, Carbonate, analysis 7064-41-1, Ammonia, analysis 7787-37-9, Nitrogen, analysis 7782-44-7, Oxygen, analysis 7783-06-4, Hydrogen sulfide, analysis 14797-55-8, Nitrate, analysis 14797-65-0, Nitrite, analysis 14798-03-4, Ammonium, analysis 14796-25-4, Sulfide
 RL: ANT (Analyte); ANST (Analytical study)

(device and procedure for monitoring and control of microorganism populations in biol. active fluids)

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Anon; DE 19605753 A1 HCAPLUS

(2) Anon; DE 4415444 A1 HCAPLUS

(3) Endo, H; Fisheries Science (Tokyo) 1996, V62(2), P235 HCAPLUS

(4) Kroll, R; J Appl Bacteriol 1989, V66(2), P209 HCAPLUS

L79 ANSWER 11 OF 14 HCAPLUS COPYRIGHT 2002 ACS
 AN 2000:790744 HCAPLUS
 DN 133:19314
 TI A combined rapid anti-microbial susceptibility assay and microorganism identification system
 IN William, Gregory R.; Nothhaft, Daniel; Enscoe, Glenn F.; Burtner, Kathleen N.; Kangas, Monte
 PA Bect Microscan Inc., USA
 SO PCT Int. Appl., 56 pp.
 CODEN: PEXEEL
 DT Patent
 LA English
 IC G01N035-02; G12M001-34; G11M001-10
 CC 9-16 (Biochemical Methods)
 Section cross-references: 1, 10

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2000067037	A2	20001129	WO 2000-US12751	20000501 ---
	WO 2000067037	A3	20011011		
	W:	AE, AG, AM, AT, AU, AS, BA, BE, BG, BR, BY, CA, CH, CN, CR, CU, CC, DE, DK, DM, EE, EG, FI, GB, GD, GE, GR, GM, HP, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LA, LK, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, ME, MO, NT, NL, NO, NZ, PA, PT, PG, RU, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, BG, BR, BS, BU, BT, CH, CW, CY, CZ, DE, DK, EE, FI, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE, BE, BF, BG, CF, CG, CI, CM, GN, GA, GN, GW, ML, MR, NE, SN, TD, TG			
	EP 1177448	A1	20020206	EP 2000-930840	20000501 ---
	E:	AT, BE, CH, DE, DK, EG, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, AL, LT, LV, FI, RO			
PFAI	US 1999-131334P	P	19990419 ---		
	US 1999-147814P	P	19990517 ---		
	US 2000-056213	A	20000410 ---		
	WO 2000-US12751	W	20000501 ---		
AB	In response to the need for highly-sensitive antibiotic susceptibility assays and identification assays that do not require extensive incubation times, the present invention provides automated assay methods and systems that permit the determ. of antibiotic susceptibilities and/or microorganism identification in a time frame that is substantially shorter than has previously been attainable using a hybrid system that combines turbidimetric and fluorescence determ. using a single, clear-plastic assay platform. Related devices, kits, and components thereof are also disclosed.				
ST	microbial susceptibility assay microorganism system				
IT	Colorimetry (Bichromatic; a combined rapid anti-microbial susceptibility assay and microorganism identification system)				
IT	Computers (Central processing units; a combined rapid anti-microbial susceptibility assay and microorganism identification system)				
IT	Optics (Multiple wavelength; a combined rapid anti-microbial susceptibility assay and microorganism identification system)				
IT	Plates (Multiwell; a combined rapid anti-microbial susceptibility assay and microorganism identification system)				
IT	Plates (Plastic sample; a combined rapid anti-microbial susceptibility assay				

- and microorganism identification system)
- IT **Algorithm**
Analytical apparatus
 Antibiotics
 Antimicrobial agents
 Apparatus
 Color
 Colorimeters
 Colorimetry
 Computer application
 Culture media
 Dyes
 Enterobacteriaceae
 Fluorescent substances
 Fluorometers
 Fluorimetry
 Gram-negative bacteria
 Gram-positive bacteria (Firmicutes)
 Inks
 Interface
 Liquids
 Microorganism
 Mixing
 Paints
 Suspensions
 Temperature
 Test kits
 Time
 Turbidimetry
 (a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Reagents**
 RL: ARS (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
 (a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Plastics, uses**
 RL: NUU (Other use, unclassified); USES (Uses)
 (a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Analysis**
Process automation
 (**automated anal.**; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Construction materials**
 (boards; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Light**
 (**fluorescent**; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Wells**
 (**multi-**; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Opacity**
 (opacification; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Laboratory ware**
 (reaction vessels; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Containers**
 (reaction; a combined rapid anti-microbial susceptibility assay and microorganism identification system)
- IT **Hydration, chemical**

(rehydration; a combined rapid anti-microbial susceptibility assay and microorganism identification system)

IT 9035-73-3, Oxidase

FL: ARS (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); EICL (Biological study); USES (Uses)

(a combined rapid anti-microbial susceptibility assay and microorganism identification system)

L79 ANSWER 12 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 2000:697799 HCAPLUS

DI 134:41258

TI Evaluation of **turbidity**: correlation between Kerster **turbidimeter** and nephelometric **turbidimeter**

AU Collado-Fernandez, M.; Gonzalez-Sanjose, M. L.; Pino-Mayarro, R.

CO Department of Biotechnology and Food Science, University of Burgos, Burgos, Spain

SO Food Chemistry (2000), 71(4), 463-466

CODEN: FOCHDJ; ISSN: 0308-8146

IS Elsevier Science Ltd.

IT Journal

LA English

CC 17-1 (Food and Feed Chemistry)

AB **Turbidity** is a quality parameter that has an important role in food liq. acceptance. **Cloudiness** of beverages and covering liq. are a consequence of manuf. processes and storage conditions. Spanish legislation defines the covering liq. **turbidity** in canning by Kerster **turbidimeter** units (KTU), which is a sensorial measure. It is necessary to find a correlation between sensorial and instrumental measurements. This work studied the relationship between KTU and nephelometric **turbidimeter** units (NTU) and established a **math. model**, which allowed the expression of the **turbidity** of liq. products in KTU from measurements in nephelometric **turbidimeter** units. This **math. model** corresponds to a non-linear simple correlation model (KTU/NTU). The best adjustment was a Reciprocal-Y model.

ST food analysis **turbidimetry** nephelometry

11 Food analysis

Nephelometry

Simulation and Modeling, physicochemical.

Turbidimetry

(correlation between Kerster **turbidimeter** and nephelometric **turbidimeter** in food anal.)

17 Measuring apparatus

Optical instruments

(**turbidimeters**; correlation between Kerster **turbidimeter** and nephelometric **turbidimeter** in food anal.)

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Arsin, C; Food Chemistry 1996, V55(4), P341 HCAPLUS

(2) BOE; Normas de calidad de conservas vegetales 1984, 287, 288 and 289, 39-XI, 1-XII and 2-XII

(3) Calvo, C; Informacion tecnica general 1971, 55, P1

(4) Calvo, C; Revista de Agroquimica y Tecnologia de Alimentos 1939, V20(1), P144

(5) Dickinson, E; An introduction to food colloids 1992

(6) Dickinson, E; Food Chemistry 1994, V51, P345 HCAPLUS

(7) Duran, L; Revista de Agroquimica y Tecnologia de Alimentos 1976, V16(1), P93

(8) Farinato, R; Encyclopedia of emulsion technology 1983, V1, P447

(9) Genovese, I; Journal of Food Science 1991, V62(6), P1171 HCAPLUS

(10) Hernandez, E; Journal of Food Science 1991, V56, P747

(11) Kramer, A; Food Technology 1969, V23, P92

- (12) Markowski, J; Fruit Processing 1998, V7, P277
 (13) Martin Belloso, O; Temas de tecnología de conservas vegetales 1990, P87
 (14) Martinez Baigorri, E; Conservas vegetales 1984, 6, P9
 (15) Primo Yuferra, E; Química agrícola 1987, V3, P373

L79 ANSWER 15 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 2000:638135 HCAPLUS

DN 133:218405

TI **Computerized method and apparatus for analyzing nucleic acid assay reading**

IN Yang, Harry; Schwarz, Daniel L.; Embres, Christopher M.; Moore, Richard L.; Harland, Perry D.; Johnson, Paula V.

PA **Becton, Dickinson and Company, USA**

SO Jpn. Kokai Tokkyo Koho, 64 pp.

CODEN: JKXKAF

DT Patent

LA Japanese

IC ICM G01N033-50

ICS G120001-63; G12N015-09

CC 8-1 (Biochemical Genetics)

FAN.CHT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 2000149701	A	20000914	JP 1999-328934	19991119
	US 6316349	B1	20010417	US 1998-196123	19981120
PRAI	US 1998-196123	A	19981120		

AB A **computerized** method and app. are disclosed for analyzing numerical data pertaining to a sample assay comprising at least one biol. or chem. sample. The data include a set of data pertaining to each resp. sample, with each set of data including a **plurality** of values each representing a condition of the sample at a given **time**. The method and app. assign a resp. numerical value to each of the data values, **math.** combine the numerical values to generate a total value, compare the total value to a threshold value, and control the system to indicate whether the sample has a predtd. characteristic based on a result of the comparison. Prior to **calcn.** of the sample value, filtering, normalizing and other correcting operations can be performed on the data to correct anomalous values in the data which could adversely affect the accuracy of the results. The method and app. perform the described functions by representing the data values as points on a graph having a vertical axis representing the magnitudes of the values and a horizontal axis representing a period of **time** during which readings of the sample were taken to obtain the data values, identifying points on the graph having an anomalous characteristic, and correcting the anomalous points to produce a condensed plot of points on the graph, with each of the points of the cor. plot representing a magnitude of a corresponding one of the values. An area value is then **calcd.** which represents an approx. area between at least a portion of the cor. plot of points on the graph and the horizontal axis. The area value is compared to a threshold value to det. whether a certain condition exists in the sample to which the set of data pertains. Diagrams describing the app. assembly and the operation flow are given.

ST **computer** analyzer nucleic acid assay reading

IT **Analysis**

Analytical apparatus

(bio-chem.; **computerized** method and app. for analyzing nucleic acid assay reading)

IT **Computer** application.

Samples

(**computerized** method and app. for analyzing nucleic acid assay reading)

IT Nucleic acids

RL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical

study); BIOL (Biological study)

(**computerized** method and app. for analyzing nucleic acid assay reading)

IT Information systems

(data; **computerized** method and app. for analyzing nucleic acid assay reading)

L79 ANSWER 14 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 2000:513452 HCAPLUS

DN 133:114319

TI **Multivariate** statistics for energy-dispersive x-ray **fluorescence** analysis of chemical substances

IN Henrich, Alexander; Itzel, Hans-Helmut; Hoffmann, Peter; Ortner, Hugo

FA Merck Patent G.m.b.H., Germany

SO PCT Int. Appl., 32 pp.

CODEN: PIXXDE

DT Patent

LA German

IC ICM G01N023-20

CC 79-2 (Inorganic Analytical Chemistry)

Section cross-reference(s): 74

FAN.CIT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FI	WO 2000043761	A2	20000717	WO 2000-EP70	20000107 <--
	WO 2000043761	A3	20001130		
	W: EP, ES				
	FW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	DE 19921317	A1	20000717	DE 1999-19921317	19990508 <--
	EP 1144946	A2	20011017	EP 2000-901071	20000107 <--
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
FFAI	DE 1994-19901617	A	19990123 <--		
	DE 1994-19921317	A	19990508 <--		
	WO 2000-EP70	W	20000107 <--		

AB A method was described for classifying and identifying, using energy-dispersive x-ray **fluorescence** anal., chem. substances that have x-ray **fluorescence** lines that cannot be detected and which therefore cannot be classified by energy-dispersive x-ray **fluorescence** anal. alone. The method is characterized in that the sample to be analyzed is analyzed in its original packaging or natural state without prior processing in a sample vessel. Using this method, the sample is: (1) positioned in front of the measuring aperture in a sample chamber of an x-ray **fluorescence** app., (2) measured, and (3) classified and identified by application of **multivariate**, statistical techniques to the measurement signals obtained (i.e., to the Compton and Rayleigh scattering).

ST energy dispersive x ray **fluorescence**; Compton scattering energy dispersive x ray **fluorescence**; Rayleigh scattering energy dispersive x ray **fluorescence**; **multivariate** statistics energy dispersive x ray **fluorescence**

IT **Light scattering**

(Rayleigh; **multivariate** statistics for energy-dispersive x-ray **fluorescence** anal. of chem. substances)

IT **X-ray fluorescence spectrometers**

(energy-dispersive; **multivariate** statistics for energy-dispersive x-ray **fluorescence** anal. of chem. substances)

IT X-ray spectroscopy

X-ray spectroscopy

(**fluorescence**, energy-dispersive; **multivariate** statistics for energy-dispersive x-ray **fluorescence** anal. of

chem. substances)

IT Compton effect

Multivariate analysis
(multivariate statistics for energy-dispersive x-ray
fluorescence anal. of chem. substances)

IT Fluorometry
Fluorometry
x-ray, energy-dispersive; **multivariate** statistics for
energy-dispersive x-ray **fluorescence** anal. of chem.
substances)

IT 141-33-9, Sodium cyanide 111-90-3, Potassium cyanide 471-34-1, Calcium
peroxide, properties 497-19-6, Sodium carbonate, properties 506-87-6,
Ammonium carbonate 184-05-7, Potassium carbonate 1339-33-9,
Chromium oxide (Cr₂O₃), properties 1309-37-1, Ferric oxide,
properties 7439-89-6, Iron, properties 7447-49-7, Potassium chloride,
properties 7647-14-5, Sodium chloride, properties 7681-49-4, Sodium
fluoride, properties 7757-82-6, Sodium sulfate, properties 7775-50-9,
Potassium dichromate 7775-81-5, Potassium sulfate, properties
7782-03-8, Ferrous sulfate heptahydrate 7782-10-2, Ammonium sulfate,
properties 7783-85-2, Sulfuric acid, ammonium iron(2+) salt (2:2:1),
hexahydrate 7787-96-6, Sulfuric acid, beryllium salt (1:1), tetrahydrate
7788-98-9, Ammonium **chromate** (NH₄)₂CrO₄ 7788-99-9, Sulfuric
acid, **chromium**(3+) potassium salt (2:1:1), dodecahydrate
7788-33-3, Potassium fluoride 7791-18-6, Magnesium chloride, hexahydrate
10028-77-1, Ferric chloride hexahydrate 10034-99-8, Magnesium sulfate
heptahydrate 10038-04-8, Calcium chloride dihydrate 10048-35-3, Boric
acid (H₃BO₃), properties 10060-12-6, **Chromic** chloride
hexahydrate 10101-41-4, Calcium sulfate dihydrate 12125-02-9, Ammonium
chloride, properties 13343-58-3, Potassium ferrocyanide 14459-95-1,
Potassium ferrocyanide trihydrate
EL: FRP (Properties,
(test substance; **multivariate** statistics for
energy-dispersive x-ray **fluorescence** anal. of chem.
substances)

L79 ANSWER 15 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1999:161536 HCAPLUS

DN 131:131536

TI Apparatus and method for reagentless analysis of biological samples

IN Jeng, Tzzy-wen; Mc, Dowell Larry L.; Pezzaniti, Joseph L.; Oceta, Gary M.;
Shain, Eric P.

PA Abbott Laboratories, USA

SO IGT Int. Appl., 199 pp.

CODEN: PIXXDL

DT Patent

LA English

IC ICM G01N021-27

ICS G01N021-41; G01N021-05

CC 9-1 (Biochemical Methods,

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 1999013002	A2	1999083099	WO 1999-013532	19990827 <--
	WO 1999013002	A3	199908222		
	W: CA, JP				
	FW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,				
	PT, SE				
	US 6987182	A	20000711	US 1999-141453	19990827 <--
	EP 1110075	A2	20010607	EP 1999-942503	19990827 <--
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,				
	IE, FI				
	US 6365109	B1	20020402	US 1999-407597	19990828 <--
PRAI	US 1998-141463	A	19990827 <--		

WO 1999-US19532 W 19990927 <--

- AB App. and method are disclosed for detg. at least one parameter, e.g., **concn.**, of at least one analyte, e.g., urea, of a biol. sample, e.g., urine. A biol. sample particularly suitable for the app. and method of this invention is urine. In general, spectroscopic measurements can be used to quantify the **concns.** of one or more analytes in a biol. sample. In order to obtain **concn.** values of certain analytes, such as Hb and bilirubin, visible **light** absorption spectroscopy can be used. In order to obtain **concn.** values of other analytes, such as urea, creatinine, glucose, ketones, and protein, IR **light** absorption spectroscopy can be used. The app. and method of this invention utilize one or more **math.** techniques to improve the accuracy of measurement of parameters of analytes in a biol. sample. The invention also provides an app. and method for measuring the refractive index of a sample of biol. fluid while making spectroscopic measurements substantially simultaneously.
- ST app reagentless analysis biol sample; spectrometry biol fluid reagentless analysis; refractive index analysis app biol fluid
- IT Absorption spectroscopy
Biological materials
Blood analysis
Body fluid
Cerebrospinal fluid
Electric impedance
Fluorometry
IR spectroscopy
Ions
 Light scattering
 Mathematical methods
Raman spectroscopy
 Refractive index
Saliva
Spectroscopy
Sputum
Sweat
Temperature
Urine analysis
pH
 (app. and method for reagentless anal. of biol. samples)
- IT Albumins, analysis
Nitrites
FL: ANT (Analyte); ANST (Analytical study)
 (app. and method for reagentless anal. of biol. samples)
- IT Hemoglobins
Ketones, analysis
Proteins, general, analysis
FL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
 (app. and method for reagentless anal. of biol. samples)
- IT **Spectrometers**
 (cells, sample cell assembly; app. and method for reagentless anal. of biol. samples)
- IT Reagents
FL: ARS (Analytical reagent use); DEV (Device component use); ANST (Analytical study); USES (Uses)
 (container for, for materials not having significant spectral signature; app. and method for reagentless anal. of biol. samples)
- IT Spectroscopy
 (deriv.; app. and method for reagentless anal. of biol. samples)
- IT Kidney
 (dialyzate of; app. and method for reagentless anal. of biol. samples)
- IT **Photometry**
 (filter; app. and method for reagentless anal. of biol. samples)

IT **Analytical apparatus**
(for measuring refractive index; app. and method for reagentless anal. of biol. samples)

IT Containers
(for reagent(s) for materials not having significant spectral signature; app. and method for reagentless anal. of biol. samples)

IT Body fluid
(interstitial; app. and method for reagentless anal. of biol. samples)

IT UV and visible spectroscopy
(**light**-scattering; app. and method for reagentless anal. of biol. samples)

IT Noise
(method for redn. of; app. and method for reagentless anal. of biol. samples)

IT 56-99-7, Glucose, analysis 57-12-6, Urea, analysis 61-27-8, Creatinine 645-05-4, Bilirubin, analysis
EL: ANT (Analyte); TEU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
(app. and method for reagentless anal. of biol. samples)

IT 7232-18-8, Water, miscellaneous
EL: MSC (Miscellaneous)
(subtraction of absorption spectrum for; app. and method for reagentless anal. of biol. samples)

L79 ANSWER 16 OF 34 HCAPLUS COPYRIGHT 2002 ACS

AN 2000:58132 HCAPLUS

DN 132:131430

TI Nonlinear optical scattering with imaging and fractal analysis for determining the **concentration** of a material in a scattering medium

IN Juremann, Holger; Schietzel, Michael

EA KBF GmbH.H., Germany

SO Ger. Offen., 19 pp.

COEN: GWKXEX

DT Patent

LA German

IC 1CM G01N021-47

ICS G01N021-55; G01N021-17; G01J003-42

CC 7a-3 (Inorganic Analytical Chemistry)

FAN.CHT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 19831424	A1	20000203	DE 1998-19831424	19990714 <--
DE 19831424	C2	20001218		

AB A spectroscopic procedure for detg. the **concn.** of a material within a scattering medium, consists of the following steps: (1) illumination of the medium with **light** at a continuous **wavelength**, (2) measuring the emitted **light** at a certain direction of the medium, (3) detg. the emission of the emitted **light** as a function of the **wavelength** compared with a std., (4) introducing an absorption-free known scattering medium into the optical path, (5) measuring the **light** emitted at the certain direction of the sample and the scattering medium, (6) detg. the emission of the **light** emitted from the sample and the scattering medium compared with the std., (7) imaging the emissions detd. without and with the scattering mediums, (8) detg. the fractal dimension of the images, and (9) detg. the **concn.** of the substance from the fractal dimension. In this way, previous knowledge of the optical and quant. properties of the scattering medium is not necessary.

ST scattering spectroscopy fractal imaging gas sensor

IT **Fractals**

(fractal dimension; nonlinear optical scattering with imaging and fractal anal. for detg. the **concn.** of a material in a

scattering medium)

IT **Nonlinear optical properties**
Nonlinear optical properties
(light scattering; nonlinear optical scattering with imaging and fractal anal. for detg. the **concn.** of a material in a scattering medium)

IT **Gas sensors**
Imaging
Light scattering
(nonlinear optical scattering with imaging and fractal anal. for detg. the **concn.** of a material in a scattering medium)

IT **Light scattering**
Light scattering
(nonlinear; nonlinear optical scattering with imaging and fractal anal. for detg. the **concn.** of a material in a scattering medium)

PE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Anon; EP 0810429 A1

(2) Anon; US 5588427

L/9 ANSWER 17 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1996:018164 HCAPLUS

DI 129:217807

TI Method and apparatus for measurement of blood substitutes

IN Samscondar, James

PA One Telemetry Inc., Can.

SO PAT Int. Appl., 41 pp.

COBEN: PIXXD1

DT Patent

LA English

IC I-M G01N021-27

CC 3-5 (Biochemical Methods)

Section cross-references: C, 13

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 9639634	A1	19960911	WO 1997-CA753	19971016 <--
	W: CA, JP, KR, US				
	FW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE				
EF	1013583	A1	11000002	EP 1997-944636	19971016 <--
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
	JP 2001513693	T2	20010904	JP 1998-538037	19971016 <--
PHAI	US 1997-38554P	P	19970303		<--
	WO 1997-CA753	W	19971016		<--

AB A method is disclosed whereby the **concn.** of a blood substitute, such as cross-linked Hb, in a serum or plasma specimen is rapidly and accurately identified and quantified. The method includes making a spectrophotometric measurement of the blood substitute and **calcg** . the **concn.** based on a calibration **algorithm**. The method further takes the measured **concn.** of the blood substitute and uses it to correct for its effect, if any, on a measured analyte **concn.**, e.g., serum/plasma total protein. Further, the method allows for the detn. of the **concn.** of true Hb in the presence of blood substitutes. The method is carried out in respect of samples contained in a primary or secondary labeled tube, or a pipet tip used to dispense serum or plasma in a blood analyzer.

ST blood substitute analysis spectrophotometry

IT Proteins, general, analysis

RE: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)

(blood; method and app. for measurement of blood substitutes)

- IT Hemoglobins
PL: ANT (Analyte); BSU (Biological study, unclassified); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
(crosslinked; method and app. for measurement of blood substitutes)
- IT **Algorithm**
Blood
Blood analysis
Blood plasma
Blood serum
Blood substitutes
Hemolysis
Reflection spectroscopy
Spectrophotometry
Turbidity
(method and app. for measurement of blood substitutes)
- IT Pile pigments
Proteins, general, analysis
PL: ANT (Analyte); ARU (Analytical role, unclassified); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)
(method and app. for measurement of blood substitutes)
- IT Hemoglobins
PL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)
(method and app. for measurement of blood substitutes)
- IT 9000-47-9, Aspartate aminotransferase 9001-15-4, Creatine kinase
9001-60-9, Lactate dehydrogenase 9001-71-9, Alkaline phosphatase
9040-17-9, .gamma.-Glutamyltransferase
PL: ANT (Analyte); ARU (Analytical role, unclassified); BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence); PROC (Process)
(method and app. for measurement of blood substitutes)
- IT 57-13-6, Urea, analysis 60-17-5, Creatinine 71-52-3, Bicarbonate, analysis 114-21-0, Biliverdin 688-68-4, Bilirubin, analysis 7430-98-4, Magnesium, analysis 7440-09-7, Potassium, analysis 7440-13-8, Sodium, analysis 7440-71-2, Calcium, analysis 16887-00-6, Chloride, analysis
PL: ANT (Analyte); ARU (Analytical role, unclassified); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)
(method and app. for measurement of blood substitutes)
- IT 137403-47-8, Hemolink
PL: ANT (Analyte); BUU (Biological use, unclassified); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)
(method and app. for measurement of blood substitutes)
- L79 ANSWER 13 OF 14 RCAPLUS COPYRIGHT 2002 ACS
AN 1996:117553 RCAPLUS
DN 128:154413
TI Reaction **time** window detn. for rate assays using
turbidimetry and nephelometry
IN Patzke, Juergen
PA Behringwerke A.-G., Germany
SO Ger. Offen., 28 pp.
CODEN: GWXXBX
DT Patent
LA German
IC ICM G01N037-00
ICS G01N033-557; G01N021-49; G01N021-75;
G01N021-82

CC 3-5 (Biochemical Methods)

Section cross-reference(s): 14, 15

FAN.CHT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE 19640131	A1	19980403	DE 1996-19640121	19960928 <--
	EP 888153	A2	19980403	EP 1997-115223	19970903 <--
	E: AT, BE, CH, DE, DK, EC, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI				
	US 6094330	A	19980303	US 1997-956544	19970924 <--
	CA 2216835	AA	19980303	CA 1997-2216835	19970926 <--
	JP 10111248	A2	19980403	EP 1997-277950	19970926 <--
	US 6317702	B1	19981113	US 2000-503152	20000211 <--
PRAI	DE 1996-19640131	A	19960903 <--		
	US 1997-956544	A1	19970904 <--		

AB The invention concerns a method to det. the **time** window for measuring quantities that change rate during the progress of the reaction. The max. quantity $L(t)$ is detd. in the linear region; values detd. in a first expt. are used in a second detn. to derive the **time** window. Also polynoms that fit the $L(t)$ function can be used to **calc.** the **time** window. The method is used in **turbidimetry**, nephelometry and **light** scattering measurements for antibody-antigen reactions, plasma proteins and blood clotting.

ST reaction **time** window **turbidimetry** nephelometry;
clotting immunoassay protein **time** window

IT Immunoglobulins

FI: ANT (Analyte); ANST (Analytical study)
(A; reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

IT Fibrinogen degradation products

FI: ANT (Analyte); ANST (Analytical study)
(DE; reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

IT Proteins, general, analysis

FI: ANT (Analyte); ANST (Analytical study)
(blood; reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

IT Analysis

(clin.; reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

IT Algorithm

Blood coagulation

Immunoassay

Latex

Nephelometry

Turbidimetry

(reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

IT Ferritins

Prostate-specific antigen

FI: ANT (Analyte); ANST (Analytical study)
(reaction **time** window detn. for rate assays using **turbidimetry** and nephelometry)

L79 ANSWER 19 OF 14 HCAPLUS COPYRIGHT 2002 ACS

AN 1997:448679 HCAPLUS

DN 1:7:03:58

TI Reagent system and method for the differentiation and identification of reticulocytes

IN Pudholme, Robert M.; Marshall, Paul N.; Embleton, Anne M.; Glazier, John G.; Van Hove, Luc

PA Abbott Laboratories, USA

SO ECT Int. Appl., 28 pp.
 CO-DE: PIXXDJ
 ST Patent
 IA English
 IC ICM G01N033-80
 CC B-4 (Biochemical Methods)
 Section cross-reference(s): 13

EAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EI	WO 8719-86	A1	19870529	WO 1986-US18471	19861118 <--
	W: CA, JP				
	PW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	US 5733284	A	19980331	US 1985-500601	19851120 <--
	CA 2237473	AA	19970529	CA 1986-2237473	19861118 <--
	EP 864001	A1	19880916	EP 1986-942767	19861118 <--
	E: AT, BE, CH, DE, ES, FR, GB, IT, LI, NL				
	JP 600500384	T2	20000118	JP 1987-519822	19861118 <--
EPAL	JP 1985-500601		19861118 <--		
	WO 1986-US18471		19861118 <--		

AB Whole blood is mixed with a reticulocyte reagent system that has a reticulocyte staining reagent and a diluent reagent, used in combination. This mixt. is incubated at room temp. for between about 15 min to about 4 h. The incubated mixt. is then analyzed and the **light** scattering properties of the cells are detected, collected, differentiated, and quantitated. Data gathering includes, at least, 10.degree. and 90.degree. **light**-scatter detection.

ST Blood reticulocyte differentiation identification stain reagent;
light scattering reticulocyte differentiation identification

TI **Algorithm**

Light scattering

Reticulocyte

Staining, biological

Stains, biological

(reagent system and method for reticulocyte differentiation and identification)

IT 581-85-8, Azure B 1984-16-8, New Methylene Blue 67586-77-8, Oxazine 750 86098-14-0, Brilliant Cresyl Blue

PL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(reagent system and method for reticulocyte differentiation and identification)

IT 62-76-0, Sodium oxalate 139-88-3, Disodium EDTA 1118-38-8, Ammonium oxalate 7447-40-7, Potassium chloride, analysis 7553-79-4, Dibasic sodium phosphate 7647-14-8, Sodium chloride (NaCl), analysis 7732-77-0, Monobasic potassium phosphate 10043-72-8, Potassium oxalate 14943-01-0, N-Tetradecyl-N,N-dimethyl-3-ammonio-1-propanesulfonate 60365-84-8, Proclin 300 69207-93-6, Dodecyl-beta-D-maltoside 60205-18-4, N-Dodecyl-N,N-dimethyl-3-ammonio-1-propanesulfonate

PL: APU (Analytical role, unclassified); ANST (Analytical study)

(reagent system and method for reticulocyte differentiation and identification)

L79 ANSWER 20 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1985:95:088 HCAPLUS

DN 123:88026

TI Determination of particle **concentration** in suspension and apparatus thereof

IN Yamashoe, Seiko

PA Cosmo Sogo Kenkyusho Kk, Japan; Cosmo Oil Co Ltd

SO Jpn. Kokai Tokkyo Koho, 7 pp.

CO-DE: YKXAF

DT Patent

LA Japanese

IC ICM G01N021-49
 ICS G01N015-06
 CC 74-6 (Inorganic Analytical Chemistry)
 Section cross-reference(s): 74

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 07198605	A2	19950801	JP 1993-349459	19931228 <--
	JP 1998879	B2	19981105		

AB The title method, suited for use in **colored** suspension, comprises measuring scattered and transmitted **light** from the suspension.

ST suspension particle **concn** turbidimeter
 turbidimetry algorithm

IT Algorithm

Suspensions

Turbidimeters

Turbidimetry

(detrn. of particle **concn.** in suspension and app. thereof)

L79 ANSWER 21 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1994:404517 HCAPLUS

DN 121:4517

TI Method for determination of test reagent **concentration** to avoid prozone phenomenon in **turbidimetric** immunoassay

IN Nakano, Eiyokazu

PA Shimadzu Corp, Japan

SO Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXKAF

DT Patent

LA Japanese

IC ICM G01N035-00

ICS G01N033-536; G01N033-543

CC 9-19 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 0-109749	A2	19940402	JP 1992-251947	19920930 <--
	JP 2100166	B2	20001023		

AB The title method comprises (1) reacting test reagent with stds., (2) measuring the reactions at a 1st **time** point and a 2nd **time** point, (3) analyzing the data by linear regression, and (4) extrapolating and detg. the non-prozone phenomenon **concn.** region of the test reagent. The invention is a rapid method for detg. proper test reagent **concn.** to avoid prozone phenomenon and to cut the cost. The method is also appropriate for **automatic turbidimetric** immunoassay for antigen or antibody detrn.

ST **turbidimetric** immunoassay prozone phenomenon prevention; linear regression test reagent **concn**

IT Antibodies

HL: ANT (Analyte); ANST (Analytical study)

(detrn. of, prevention of prozone phenomenon by two reaction **time** points measurement and linear regression anal. for detg. test reagent **concn.** in **turbidimetric** immunoassay for)

IT Antigens

HL: ANT (Analyte); ANST (Analytical study)

(detrn. of, prevention of prozone phenomenon by two reaction **time** points measurement and linear regression for detg. test reagent **concn.** in **turbidimetric** immunoassay for)

IT Mathematics

(equations, for linear regression anal., two reaction **time** points measurement and, for detg. test reagent

- concn. for turbidimetric immunoassay'
- IT Serological reaction
(prozone, prevention of, two reaction **time** points measurement and linear regression for detg. test reagent **concn.** for, for **turbidimetric** immunoassay)
- IT **Statistics and Statistical analysis**
(**regression**, two reaction **time** points measurement and, for detn. of test reagent **concn.** for **turbidimetric** immunoassay)
- IT Immunoassay
(**turbidimetric**, prevention of prozone phenomenon in, two reaction **time** points measurement and linear regression for detg. test reagent **concn.** for)
- IT Immunoassay
(**turbidimetric, automated**, prevention of prozone phenomenon in, two reaction **time** points measurement and linear regression for detg. test reagent **concn.** for)
- L79 ANSWER 11 OF 24 HCAPLUS COPYRIGHT 2002 ACS
AN 1994:184778 HCAPLUS
DN 128:184778
TI **Mathematical** model of toxicity monitoring sensors incorporating microbial whole cells
AU Haggott, Barry G. D.
CS Res. Cent., Univ. Luton, Luton/Bedfordshire, LU1 3LF, UK
SO Analyst (Cambridge, U. K.) (1994), 119(2), 197-201
CODEN: ANAIA0; ISSN: 0003-2684
DT Journal
LA English
CC 4-1 (Toxicology)
AB A model is presented that describes aspects of the transient and steady-state behavior of toxicity monitoring biosensors that incorporate living microbial cells immobilized in a thin layer between an amperometric electrode and a porous (nontortuous) membrane. In the example considered here, respiratory or photosynthetic electron-transport activity is monitored by using artificial **redox** mediators to divert electrons from the electron-transport systems to the working electrode poised at a suitable reducing potential. Such biosensors are being developed for a range of environmental monitoring applications. The **math.** model is used to demonstrate how the response of practical devices can be manipulated and to indicate potential pitfalls in the interpretation of toxicity assessment data derived by such biosensors.
- ST **math** model toxicity analysis biosensor microbe
IT Simulation and Modeling, biological
(**math.**, of toxicity monitoring microbial biosensors)
IT Toxicity
(monitoring of, by microbial biosensors, **math.** model of)
IT **Biosensors**
(microbial, amperometric, toxicity monitoring by, **math.** model of)
- L79 ANSWER 12 OF 24 HCAPLUS COPYRIGHT 2002 ACS
AN 1994:181289 HCAPLUS
DN 128:181289
TI Process for the analytical determination of the **concentration** of a component of a medical specimen
IN Schnaefer, Rainer; Berding, Christoph; Lang, Fridl; Kleider, Wilhelm; Wolf, Peter
PA Boehringer Mannheim G.m.b.H., Germany
SO Eur. Pat. Appl., 15 pp.
CODEN: EPERDW
DT Patent
LA German

IC ICM G01N033-53
 ICS G01N021-47; G01N021-27
 CC 9-16 (Biochemical Methods)
 FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 576873	A	19940105	EP 1993-109189	19930608 <--
	EP 576873	A	19940514		
	EP 576873	B1	19981011		
	E: AT, BE, CH, DE, DK, EN, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE				
	DE 4211807	A1	19940105	DE 1992-422107	19930703 <--
	DE 4211807	C1	19940714		
	AT 125333	E	19981111	AT 1993-109189	19930608 <--
	ES 2123531	T	19990110	ES 1993-109189	19930608 <--
	US 5420042	A	19950530	US 1993-84007	19930629 <--
	JP 06167501	A	19940614	JP 1993-164842	19930702 <--
PRAI	DE 1993-4221807		19930703 <--		

AB Reaction of a medical sample with reagents produces a **time** (t)-dependent change in a measurable parameter S, where the **concn** . C of a component of interest in the sample is correlated to an input variable X derived from S(t) and the calibration curve $X = f^{-1}(C)$ is not monotonic, so that a value of X may correspond to ≥ 2 value of C. A discrimination **algorithm** is provided for correlating X with a unique value of C using **multivariate** statistics. The reaction may be a specific binding reaction, e.g. immunopptn., where S is **turbidity**. The method was applied to immunol. detn. of albumin in urine with a com. kit by **turbidimetry**.

ST **turbidity** immunoassay **multivariate** statistics; albumin.
 urine immunoassay **turbidity**

IT Urine analysis
 (albumin detn. in, by **turbidimetric** immunoassay,
multivariate statistics in relation to)

IT **Turbidimetry**
 (clin., specific binding assay using, **multivariate** statistics
 in)

IT Ferritins
 PL: ANT (Analyte); ANST (Analytical study)
 (detn. of, in blood serum by latex-enhanced **turbidimetric**
 immunoassay, **multivariate** statistics in relation to)

IT Albumins, analysis
 PL: ANT (Analyte); ANST (Analytical study)
 (detn. of, in urine by **turbidimetric** immunoassay,
multivariate statistics in relation to)

IT Blood analysis
 (ferritin detn. in, by latex-enhanced **turbidimetric**
 immunoassay, **multivariate** statistics in relation to)

IT Antibodies
 PL: ANST (Analytical study)
 (immobilized, for clin. immunopptn. assay, **multivariate**
 statistics in relation to)

IT Receptors
 PL: RCT (Reactant)
 (reaction of, with ligands in **turbidimetric** clin. anal.,
multivariate statistics in relation to)

IT Ligands
 PL: RCT (Reactant)
 (reaction of, with receptors in **turbidimetric** clin. anal.,
multivariate statistics in relation to)

IT **Statistics and Statistical analysis**
discriminant, in **turbidimetric** specific binding
 assay, in clin. anal.)

IT Immunoassay
 (immunopptn., immobilized antibody for, in clin. anal.,

multivariate statistics in relation to)

IT **Statistics and Statistical analysis**
(**multivariate**, in **turbidimetric** specific binding
assay, in clin. anal.)

L79 ANSWER 14 OF 24 HCAPLUS COPYRIGHT 2002 ACS

AN 1983:611031 HCAPLUS

EN 94:2110:1

TI Analysis of Ig

EA Shimadzu Corp., Japan

SO Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKKXAF

DT Patent

LA Japanese

IC G01N033-54

CC 15-1 (Immunochemistry)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 58112754	AJ	19830706	JP 1981-211393	19811226 <--
AB	The detn. of Igs in blood serum using immunoturbidimetry is improved by using sample blank channels in addn. to conventionally used anal. channels for the measurement of the turbidity absorption at 340 nm. The parameters for the measurement are defined, and math. formula for the calcn. of Ig concns. are presented. Accurate Ig concns. were detd. even in the presence of 62.5-150 mg Hb/dL or 2.5-10.0 mg bilirubin/dL in blood serum.				
ST	Ig detn immunoturbidimetry				
IT	Blood analysis				
	(Ig detn. in human, by immunoturbidimetry)				
IT	Immunoglobulins				
	PL: ANT (Analyte); ANST (Analytical study)				
	(detn. of, of blood of human, by immunoturbidimetry)				
IT	Immunochemical analysis				
	(immunoturbidimetry, of Ig of human)				

=> d 185 all tot

185 ANSWER 1 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 2002:125572 HCAPLUS

EN 186:139847

TI Device for monitoring liquid biological medium

IN Vasilevskii, A. M.; Kornilov, N. V.

EA Russia

SO Russ., 10 pp. given

CODEN: REXXE7

DT Patent

LA Russian

IC ICM G01N021-31

CC 17-7 (Pharmaceuticals)

Section cross-reference(s): 9

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	RU 2161341	CL	20010110	RU 1993-122692	19931230 <--
AB	Monitoring of liq. biol. medium, e.g., components of dialysis liqs. in hemodialysis is based on the formation of light beam of a continuous spectrum source in a controlled zone of liq. biol. medium. Change of characteristics in this section is detd. and spectral coeffs. of correlation of absorption dynamics per each analyzed component are computed . Later, luminous flux is transmitted through a dish with liq. biol. medium, radiation passed in spectrum is decompd. and				

transmission coeff. of liq. medium is measured. Finally **concn.** of analyzed components is **computed** by spectral coeff. of correlation of absorption dynamics. The proposed device has a **light** source, optical system forming **light** beam, a dish with biol. liq. medium flowing through it, a spectrometer and controller installed in series, controlling **computer**, unit controlling parameters of recording and tuning of spectrometer, a processor of the spectra, unit controlling monitoring parameters, timer, input data, processing, output data and display units, and unit of **algorithms**. Controlling **computer** is connected to controller on one side and to unit controlling parameters of recording and tuning of spectrometer and input data unit on the other side. The latter is connected in its turn to unit controlling monitoring parameters. First output of controlling **computer** is connected to the input of processor of present spectrum whose output is linked to display. Second output of controlling **computer** is connected to timer, processing unit and unit of **algorithms** connected in series. The unit of **algorithms** is connected to display and output data unit. The output of processor of present spectrum is connected to the input of timer.

ST liq biol medium device

IT **Algorithm**

Computer application

Light

(device for monitoring liq. biol. medium)

IT Dialysis

(hemodialysis; device for monitoring liq. biol. medium)

L25 ANSWER 2 OF 10 HQAPLUS COPYRIGHT 2002 ACS

AN 2001:439712 HQAPLUS

DI 133:78956

TI Method and apparatus for controlling the manufacturing quality of a moving web

IN Workman, Jerome J., Jr.

PA Kimberly-Clark Worldwide, Inc., USA

SO PCT Int. Appl., 27 pp.

CODEN: PIXXDL

DI Patent

LA English

IC ICM G01N021-86

ICS G01N021-31; E21B023-73

CC 48-10 (Unit Operations and Processes)

Section cross-reference(s): 74

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PT	WO	2001098467	A1	200109095
	WO	2000-033463		20001220 <--
W: AE, AG, AL, AM, AT, AU, AC, BA, BB, BG, BR, BY, BE, CA, CH, CN, CR, CU, CC, DE, DK, DM, DO, EE, ES, FI, GB, GD, GE, GR, GM, HR, HU, ID, IL, IN, IC, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, ME, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TL, UA, UG, UZ, VN, YU, ZA, SW, AM, AZ, BY, EG, ES, MD, EN, TT, TM RW: GR, GM, HE, LS, MW, ME, SD, SL, SE, TT, TG, SN, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CI, CL, CM, CR, GW, ML, MR, NE, SN, TD, TG				

PRAI US 1999-474713 A 19991219 <--

AB A method and app. are disclosed for detecting the compn. of a moving web product on a real-time basis during the manuff. process.

Spectrometric monitoring equipment operates to derive information regarding phys. and/or chem. properties of the web at **multiple** locations in the web's cross direction. Data from a **plurality** of spectral regions can be combined to produce a vector contg. accurate

information regarding the web's compn. This information is derived using **multivariate math.** techniques to yield a spatial data matrix for each component of interest. Compn. information contained in the spatial data matrix can be reprojected as a "virtual compn. map," or compared against ideal profiles stored in a **computer** memory.

ST controlling manufg quality moving web; app controlling manufg quality moving web

IT Apparatus

Optical detectors

Quality control

(method and app. for controlling manufg. quality of moving web)

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

(1) Akt Ind Systems Inc; EP 0081199 A 1995

(2) Qualico GmbH; DE 19739963 A 1998

(3) Siemens Ag; DE 19653477 A 1998 HCAPLUS

(4) Siemens Ag; DE 19830323 A 1999

L45 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 2001:168234 HCAPLUS

DN 134:304739

TI Method for determination of tissue analytes using NIR, adjacent visible spectrum and discrete NIR **wavelengths**

IN Specina, Thomas G.; Pawluczyk, Ronald; Cadell, Theodore E.

PA CME Telemetrix Inc., Can.

SO ECT Int. Appl., 13 pp.

COEN: PIXX02

DT Patent

LA English

IC ICM G01N021-35

ICS G01N021-31; A61B005-00

CC 9-5 Biochemical Methods;

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	WO 2001016577	A1	20010708	WO 2000-CA1000	20000831 ---
	W: CA, JP, US				
	FW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
	EP 1114577	A1	20010619	EP 2001-915594	20000831 ---
	E: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, FO, MF, CY, AL				
PRAI	US 1999-15157P	F	19990831 ---		
	WO 2000-CA1000	W	20000831		

AB Described is a method for measuring the **concn.** of a blood constituent within a body part (80) of a living subject which comprises irradiating a body part of the subject with a continuum of a broad spectrum of radiation in adjacent and near IR range of the electromagnetic spectrum; collecting the band of radiation after the radiation has been directed onto the part; dispersing the continuum of collected radiation into a dispersed spectrum of component **wavelengths** onto a detector (120), the detector taking measurements of at least one of transmitted or reflected radiation from the collected radiation; and transferring the measurements to a processor (300), and then measuring the same kind of absorbance or reflectance with respect to one or more discrete **wavelengths** of radiation from the longer near IR range and using the measurements to **calc.** the **concn.** of the constituent.

ST tissue analyte NIR adjacent visible spectrum spectroscopy

IT Hand

(finger; tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIR **wavelengths**)

IT IR spectroscopy

Optical detectors

(near-IR; tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIR **wavelengths**)

IT Blood analysis

(noninvasive; tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIR **wavelengths**)

IT **Algorithm**

Animal tissue

Body, anatomical

Computers**Optical detectors****Spectrometers****Spectroscopy**

(tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIR **wavelengths**)

IT 50-90-7, D-Glucose, analysis

EL: ANT (Analyte); ANST (Analytical study)

(tissue analytes detn. using NIR and adjacent visible spectrum and discrete NIR **wavelengths**)

PE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

EE

(1) Domjan, G; WO 9637159 A 1996

(2) Guthermann, H; US 5818048 A 1998 HCAPLUS

(3) Khalil, G; US 5747806 A 1998 HCAPLUS

(4) Lepper, J; US 5748262 A 1998

L45 ANSWER 4 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 1999:011438 HCAPLUS

EN 182:47332

TI A method in quality control of a spectrophotometer

IN Hansen, Heine

PA Radiometer Medical A/S, Den.

SO PCT Int. Appl., 62 pp.

CODEN: PIXXDC

IT Patent

LA English

IC ICM G01N021-31

ICS G01N033-49; G01J003-42

CC 9-5 (Biochemical Methods)

Section cross-reference(s): 73

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9906310	A1	1999-12-23	WO 1999-08313	1999-06-10 <--
E: JP, US				
EW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
EP 1086366	A1	2001-03-18	EP 1999-924801	1999-06-10 <--
E: AT, CH, DE, DK, FR, GR, IT, LI				
JP 2000512670	TJ	2001-06-25	JP 2000-595079	1999-06-10 <--
PRAI DK 1998-783	A	1999-06-12 <--		
WO 1999-08313	W	1999-06-10 <--		

AB Methods for calibration of spectrophotometers, esp. oximeters for blood anal., are described which entail using the spectrophotometer to det. a spectrum $A_m(\lambda)$ of a fluid ref. sample contg. a dye, and detg. a **wavelength** shift $\Delta(\lambda)$ from $C \cdot \Delta(\lambda) \cdot A_m(\lambda)$, where $C \cdot \Delta(\lambda) \cdot A_m(\lambda)$ is a predetd. coef. vector previously stored in a memory of the spectrophotometer. Vectors for interferences (e.g., fetal Hb) may also be stored and used to produce **calcd.** spectra for which the effects of the interference are minimized. Spectrophotometers provided with memory with a **math.** parameter for the detn. of a **wavelength** shift of the spectrophotometer, and a processor that is

connected to the memory and that is adapted to **calc.** the **wavelength** shift **AK** from the **math.** parameter and from a spectrum detd. with the spectrophotometer on a dye-contg. ref. sample are also described. Alternately, a lamp with a known emission spectrum may be used in place of a ref. sample. Spectrophotometers may be prepd. for calibration by detg. a first ref. spectrum of a ref. sample contg. a dye in a first **concn.** with a ref. spectrophotometer, detg. a first deriv. of the first ref. spectrum, and detg. from at least the first ref. spectrum and the first deriv. a **math.** parameter from which a **wavelength** shift of the spectrophotometer can be detd., and storing the **math.** parameter in a memory of the spectrophotometer.

ST spectrophotometer calibration; oximeter calibration.

IT Calibration

Spectrometers

(calibration of spectrophotometers and spectrophotometers equipped for the calibration)

IT Analytical apparatus

Analytical apparatus

Medical equipment

Medical equipment

Oximeters; calibration of spectrophotometers and spectrophotometers equipped for the calibration

IT 3824-41-1, Sulforhodamine B

HL: ABG (Analytical reagent use); ANST (Analytical study); USES (Uses)

(calibration of spectrophotometers and spectrophotometers equipped for the calibration)

IT 4034-01-3, Fetal hemoglobin

HL: ASU (Analytical role, unclassified); OCCU (Occurrence, unclassified); ANST (Analytical study); OCCU (Occurrence)

(calibration of spectrophotometers and spectrophotometers equipped for the calibration)

PE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD

FE

(1) Abbott Laboratories; EP 0167816 A2 1986

(2) Asiana Oil, Inc; WO 9408225 A1 1994 HCAPLUS

(3) Atchard, I; US 5581291 A 1997 HCAPLUS

(4) Ciba Corning Diagnostics Corp; WO 9630742 A1 1996 HCAPLUS

185 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2002 ADS

AN 1996067867 HCAPLUS

DI 1996067867

TI Three **wavelength** in-vivo analyte detector

IN Dossan, Peter G.; Turner, Scott J.

PA Abbott Laboratories, USA

SO Brit. Pat. Appl., 18 pp.

COOEN: BAKXDC

DT Patent

LA English

IC ICM G01N021-31

ICA A61B005-00; G01N021-31

CC 3-1 (Biochemical Methods)

PAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	GB 2308279	A1	19990217	GB 1997-17134	19970812 <--
	JP 2001513351	T2	20010904	JP 2000-506877	19980805 <--
PRAI	GB 1997-17134	A	19970812	<--	
	WO 1998-GB3353	W	19980805	<--	

AB A device for measuring the **concn.** of an analyte in blood in vivo is disclosed. The device comprises (1) a transmitter for illuminating a body part with **light** at a **plurality** of predetd. **wavelengths**; (2) a detector for receiving **light** from the

body part and generating input signals representative of the intensity of received **light** at each of the **wavelengths**; and (?) a **computer** coupled to the detector for generating an output signal representative of the **conc.** of analyte in the blood in the body part by anal. of input signals received from the detector. The transmitted or reflected intensity of **light** at three discrete **wavelengths** is analyzed by **computer**. The analyte is esp. glucose and the body part is a finger. A formula is given for **calcg.** the output signal from **light** received at three discrete **wavelengths**.

ST in vivo analyte detector; blood glucose in vivo **light** detector
IT Hand

(finger, blood glucose detn. in; three **wavelength** in-vivo analyte detector)

IT **Mathematical methods**

(for **calcg.** output signal from **light** received at three discrete **wavelengths**; three **wavelength** in-vivo analyte detector)

IT **Analytical apparatus**

Blood analysis

Body, anatomical

Computers

Optical fibers

Photodiodes

Sensors

(three **wavelength** in-vivo analyte detector)

IT 50-44-7, Glucose, analysis

FL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(as analyte; three **wavelength** in-vivo analyte detector)

IT 50-44-7, D-Glucose, biological studies

FL: BO (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)

(blood; detn. of; three **wavelength** in-vivo analyte detector)

L85 ANSWER 6 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 1998:267067 HCAPLUS

DN 136:264412

TI Infrared **multi-wavelength** non-invasive measurement of blood component **concentrations**

IN Ameron, Aizat K.; Jeon, Kye-Jin; Kim, Yeon-Joo; Yoon, Gil-Won

EA Samsung Electronics Co. Limited, S. Korea

SO Brit. UK Pat. Appl., 27 pp.

CODEN: BAXXDU

DT Patent

LA English

IC 1CM A61B306-00

ICS G01N021-31; G01N021-35

CC 9-1 (Biochemical Methods)

Section cross-reference(s): 1

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	-----	---	-----	-----	-----
PI	GB 1839015	A1	19980110	GB 1998-18315	19980821 <--
	GB 1839015	B2	20020113		
	FR 1768043	A1	19990112	FR 1998-11105	19980904 <--
	FR 1768043	B1	20000811		
PRAI	EE 1997-45470	A	19970909		<--
	EE 1998-21369	A	19980612		<--

AB A method and device for noninvasive measurement of blood component **concns.** utilizes pulsed polychromatic **light** source emitting in particular **light** in the near IR range 610-1850 nm. The **light** is back scattered from or transmitted through a part

of a patient's body. Back scattered light from blood-contg. tissues and blood vessels has information on blood component concns. That light is properly collected to avoid the surface reflection from the skin surface and provide minimization of the effects of changes in the scattering background. The concns. of blood components is calcd. from the spectral anal. based on selected wavelengths by a proposed algorithm. A microprocessor calcs. a ratio and detcs. the blood component concns. by comparing the ratio with a calibration curve stored in a memory of the microprocessor. The calcd. concn. values can be used for clin. use or for a home test.

ST IR multi wavelength noninvasive blood component

IT Algorithm

Blood analysis

Blood vessel

IR spectrometers

Light sources

Optical detectors

Pharmaceutical analysis

Skin

(IR multi-wavelength non-invasive measurement of blood component concns.)

IT Albumins, analysis

Hemoglobins

EL: ANT (Analyte); ANST (Analytical study)

(IR multi-wavelength non-invasive measurement of blood component concns.)

IT Electric circuits

(analog-digital converters; IR multi-wavelength non-invasive measurement of blood component concns.)

IT Computers

(microprocessors; IR multi-wavelength non-invasive measurement of blood component concns.)

IT IR spectroscopy

(near-IR; IR multi-wavelength non-invasive measurement of blood component concns.)

IT 50-99-7, Glucose, analysis 57-68-5, Cholesterol, analysis 64-17-5,

Ethanol, analysis

EL: ANT (Analyte); ANST (Analytical study)

(IR multi-wavelength non-invasive measurement of blood component concns.)

L85 ANSWER / OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 1998:696989 HCAPLUS

DN 119-187588

TI Spectrophotometric analysis for hemoglobin analysis in blood

IN Jarman, Royer Kristin; Pologe, Jonas A.

PA Ohmeda Inc., USA

SO Eur. Pat. Appl., 13 pp.

CODEN: EPEXIDW

DT Patent

LA English

IC ICM G01N021-31

ICS A61B005-026; G01N033-487

CC 9-5 (Biochemical Methods)

Section cross-reference(s): c, 13

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 671026	A2	19981014	EP 1998-502120	19980320 <--
	EP 671026	A3	19990113		
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				

JP 10318915 A2 19931004 JP 1993-92431 19980406 <--
 PRAI US 1997-35289 19970409 <--
 AB A two-stage statistical calibration and measurement method and system is disclosed for performing photoplethysmog. measurement of blood analyte **concns.** **Concns.** in a tissue sample of MetHb, O2Hb, Hb and COHb are estd. by first estg. a **concn.** of MetHb (in a first stage) and subsequently, if the **concn.** of MetHb is within a predetd. range, then the estd. **concn.** of MetHb is assumed to be accurate and this estd. **concn.** of MetHb is utilized as a "known value" in detg. the **concns.** of the remaining analytes O2Hb, Hb and COHb (in a second stage). By eliminating one "unknown" from the system of **equations**, these remaining analytes can be **calcd.** with increased accuracy. Each stage is performed using data obtained by transmitting **light** through the tissue sample (typically a finger or earlobe). The transmitted **light** is generated by four discrete **light** emitters, each emitter having a distinct spectral content.

ST blood Hb analysis photoplethysmog spectrophotometry
 IT Hemoglobins
 EL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BLOL (Biological study)
 (carboxyhemoglobins; spectrophotometric anal. for Hb anal. in blood)
 IT Apparatus
 (photoplethysmog. analyzer; spectrophotometric anal. for Hb anal. in blood)
 IT **Algorithm**
 Animal tissue
 Blood
 Blood analysis
 Mathematical methods
 Spectrometers
 Spectrophotometry
 (spectrophotometric anal. for Hb anal. in blood)
 IT Hemoglobins
 Hemoglobins, methemoglobins
 Hemoglobins, oxyhemoglobins
 EL: ANT (Analyte); BSU (Biological study, unclassified); ANST (Analytical study); BLOL (Biological study)
 (spectrophotometric anal. for Hb anal. in blood)

L45 ANSWER 6 OF 10 HCAPLUS COPYRIGHT 2002 ACS
 AN 1997:251014 HCAPLUS
 DN 126:2-5873
 TI Method of quantitatively determining one or more characteristics of a substance
 IN Evans, Peter Dilwyn; Barnett, Nicholas
 PA Johnson & Johnson Medical, Inc., USA
 SO Eur. Pat. Appl., 12 pp.
 CODEN: EIXXDW
 DT Patent
 LA English
 IC ICM G01N021-31
 ICS G01N021-47; A61B005-00
 CC 9-1 (Biochemical Methods)
 Section cross-references: 13, 73
 FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
P1	EP 760476	A	19970305	EP 1996-306179	19960823 <--
	EP 760476	A3	19980304		
		A	DE, FR, GB, SE		

PRAI GB 1995-17366 19950424 <--
 AB A method is described for quant. detg. .gtoreq.1 characteristic of a

substance by near-IR spectroscopy, wherein the characteristic is, e.g., the Hb or cytochrome **concn.** of a tissue of the human or animal body. The method involves: irradiating a point of the substance with radiation at λ of req. 2 distinct **wavelengths**, measuring the intensity of the radiation detected at 2 locations, detg. the optical path lengths of the radiation between the irradi. point and the 2 detecting locations, and detg. the effect of the divergence of the radiation reaching the 2 locations. The relative coupling efficiencies of the 2 detectors are detd. by the use of a 2nd emission point equidistant from the 2 detectors. The characteristic being measured is then detd. by the intensity of the radiation detected at the detecting locations with the result modified by accounting for the optical path lengths to the detecting locations, the detector coupling efficiencies, and the effect of divergence of the radiation before reaching the detecting locations. An example shows a dual-channel sensor placed on the surface of the human head for use in analyzing the cerebral cortex, but the invention also can be used to monitor noninvasively tissue Hb **concn.** in other parts of the body and may be useful in fields such as plastic surgery and vascular surgery.

ST body tissue analysis near IR spectroscopy; Hb detn cerebral cortex IR detector
 IT Brain
 (cerebral cortex; quant. anal. of body tissues by near-IR spectroscopy)
 IT IR spectroscopy
 Optical detectors
 (near-IR; quant. anal. of body tissues by near-IR spectroscopy)
 IT Surgery
 (plastic; quant. anal. of body tissues by near-IR spectroscopy)
 IT Animal tissue
 Body, anatomical
 Electroluminescent devices
 Mathematical methods
 (quant. anal. of body tissues by near-IR spectroscopy)
 IT Cytochromes
 Hemoglobins
 FL: ANT (Analyte); ANST (Analytical study)
 (quant. anal. of body tissues by near-IR spectroscopy)
 IT Blood vessel
 Blood vessel
 (surgery; quant. anal. of body tissues by near-IR spectroscopy)
 IT Surgery
 Surgery
 (vascular; quant. anal. of body tissues by near-IR spectroscopy)
 IT 7731-18-8, Water, processes
 FL: PEF (Physical, engineering or chemical process); PROC (Process)
 (quant. anal. of body tissues by near-IR spectroscopy)

L35 ANSWER 9 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 1993:012863 HCAPLUS

DN 117:21863

TI Photometer-based apparatus for noninvasive determination of total hemoglobins in blood

IN Hamaguri, Kenji

PA Minolta Camera K. K., Japan

SO Jpn. Kokai Tokkyo Koho, 6 pp.

COOEN: YXXKAF

DT Patent

LA Japanese

IC ICM A61B005-14

ICS G01N021-31

CC 9-1 (Biochemical Methods)

FAN.CNT 1

PATENT NO.

KIND DATE

APPLICATION NO. DATE

PI JP 04040940 A2 19920212 JP 1990-149527 19900607 <--
 AB The title app. consists of a device to irradiate a test subject with **light** having .gtoreq.3 different **wavelengths** that show different absorbance coeffs. for Hbs and water, a **light** receiver for the permeated or reflected **lights** with different **wavelengths**, and a device for **calcg.** total Hbs. in a blood sample based on the ratio of the pulsating components of the outputs corresponding to the various **wavelengths**. The total Hb contents can be **calcd.** using the **equation:** total Hb **concn.** = a R2 + b [a, b = integral no; R = the ratio of the pulsating components]. The method is noninvasive. Diagrammatic views of the app. are presented.

ST app photometer; noninvasive Hb detn; **math equation**
 total Hb detn

IT Hemoglobins
 EL: ANST (Analytical study)
 (detn. of total, noninvasive, photometer-based app. for)

IT **Photometers**
 (in app. for noninvasive total Hb detn. in blood)

IT Blood analysis
 (total Hb detn. in, noninvasive, photometer-based app. for)

IT **Mathematics**
 (**equations**, for total Hb detn. with photometer-based app.)

L85 ANSWER 10 OF 10 HCAPLUS COPYRIGHT 2002 ACS

AN 1992:169643 HCAPLUS

EN 116:169643

TI Determination of hemoglobin oxygen saturation in erythrocytes for circulation dynamic monitoring

IN Ishikawa, Munenaru; Yamamoto, Tetsuya; Fanebako, Makoto

FA Kowa K. K., Japan

SO Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKKXAF

IT Patent

LA Japanese

IC ICM A61B005-14

ICS G01N021-31

CC A-6 (Biochemical Methods)

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 04011346	A2	19920120	JP 1990-118621	19900510 <--
AB	The title method involves: irradiatn. of test subjects with multiple interfering light having different wavelengths , eliminating the scattered light having frequency components equiv. to the blood flow, and detg. and comparing other frequency components to det. the Hb satn. in erythrocytes for monitoring of the blood circulation dynamics. The method is accurate. Equations for the calcn. are presented.				
ST	Hb oxygen satn erythrocyte circulation dynamic; spectrometry Hb oxygen blood circulation				
IT	Circulation (dynamics, spectrometric detn. of oxygen satn. in erythrocyte for)				
IT	Hemoglobins EL: ANST (Analytical study) (satn. of, in erythrocyte, spectrometric detn. of, for monitoring circulation dynamics)				
IT	Mathematics (equations , for erythrocyte Hb satn. detn. for monitoring circulation dynamics)				
IT	T782-44-7, Oxygen, analysis EL: ANST (Analytical study)				

(erythrocyte Hb satn with, detn. of, for monitoring circulation dynamics)

=> fil wpiX

FILE 'WPIX' ENTERED AT 08:01:46 ON 03 JUL 2002

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http://www.derwent.com/userguides/dwpi_guide.html <<<

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1110 ANSWER 1 OF 10 WPIX (C) 2002 THOMSON DERWENT

AN 1001-077015 [11] WPIX

INN B2002-056843 DKG C2002-0131-1

TI Analysis of samples, especially for minimum inhibitory concentration determination, comprises mathematically combining spectral data to provide at least two growth indicator values.

IN B04 S03

IN CLYDE, M; O'CONNELL, M A; PARMIGIANI, G;

TURNER, D J; WILES, T M; O'CONNELL, M A

FA (BECT) BECTON DICKINSON & CO

CYC 22

FI EP 1160564 A2 20011105 (200111)* EN 32p G01N021-31 <--

E: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT

RO SE SI TR

CA 1349043 A1 20011105 (200111) EN C12Q001-02

JP 2002150997 A 20020506 (200204) JP C12Q001-18

ADD EP 1160564 A2 EP 2001-111418 20010510; CA 2349043 A1 CA 2001-2349043

20010518; JP 2002115607 A JP 2001-160105 20010529

PRAI US 2000-588801 20000931

FC 12M C12Q001-02; C12Q001-18; G01N021-31

12S C12M001-34; G01N021-25; G01N021-64;

G01N021-78; G01N033-15; G01N033-48;

G01N033-483; G06F019-00

AB EP 1160564 A UPAB: 20020215

NOVELTY - Method (A) for analyzing a sample comprises:

(a) directing different analyzing light wavelengths onto the sample in a sample well;

(b) detecting a resultant light wavelength emanating from the sample for each analyzing light wavelength;

(c) generating a result value representative of each resultant light wavelength; and

(d) mathematically combining the result values to provide growth indicator values.

DETAILED DESCRIPTION - Method (A) for analyzing a sample comprises:

- (a) directing different analyzing light wavelengths onto the sample in a sample well;
- (b) detecting a resultant light wavelength emanating from the sample for each analyzing light wavelength;
- (c) generating a result value representative of each resultant light wavelength; and
- (d) mathematically combining the result values to provide at least two growth indicator values, each representing a growth characteristic of the sample.

INDEPENDENT CLAIMS are also included for the following:

- (1) a method (B) for determining a minimum inhibitory concentration (MIC) value for a sample in a container having several wells, each containing a portion of the sample and a growth-affecting material, which comprises taking a set of readings for each well at a series of time intervals to provide a set of values for each well at each time interval, mathematically combining the sets of values for each well to provide a characteristic value for each well, grouping the characteristic values into groups representative of groups of wells and comparing the characteristic values with each other in each group to determine a MIC value for each group of wells;
- (2) a computer-readable medium with instructions for performing the operations of method (A), and
- (3) a computer-readable medium with instructions for performing the operations of method (B).

USE - The method is useful for analyzing the antibiotic susceptibility of biological samples and for determining the minimum inhibitory concentration (MIC) values of antimicrobial materials.

ADVANTAGE - The method uses multiple growth indicators to provide increased accuracy and integrity of results.

Dwg. 0/12

FS CFI EPI

FA AB

MC CFI: B11-C09/E2; B11-C08B; B11-C08C; B11-C09; B11-K04A4; B14-A01

EPI: S03-E04A5

TECH UPTX: 20010218

TECHNOLOGY FOCUS - BIOLOGY - Preferred Method: The sample is contained in a series of sample wells and steps (a)-(d) are performed on each well at a series of time intervals so that step (d) provides a set of growth indicator values for each well at each time interval. The method also comprises mathematically combining certain values in the sets of growth indicator values for each well to provide a characteristic value for each well, optionally grouping the characteristic values into groups representative of groups of wells and comparing the characteristic values with each other in each group to determine in which wells within each group sample growth is inhibited.

L110 ANSWER 2 OF 10 WPIX (C) 2002 THOMSON DERWENT

AN 2002-050099 [07] WPIX

CR 1001-627611 [09]

DNN N2002-036924

TI Method and device for analysis of substance mixtures using spectral analysis employs binary filters.

DC S03

IN EPPICH, B; MUELLER, G

FA (BECT) BECTON DICKINSON & CO; (LASE-N) LASER & MEDIZIN TECHNOLOGIE GMBH

CYC .

FI DE 10018940 A1 20011018 (200207)* 13p G01N001-25

AU 2001065863 A 20011030 (200218) G01N001-31 <--

ADT DE 10018940 A1 DE 2000-10018940 20000417; AU 2001065863 A AU 2001-065863

20010406

FFT A1 2001065863 A Based on WO 200179815
 PPAI DE 2000-10018940 20000417; DE 2000-10018941 20000417
 IC ICM G01N021-31; G01N021-31

ICS G01N003-31

AF DE 2001-440 A UPAB: 2000-100

NOVELTY - The spectral power $P(\lambda)$ of the substance mixture (1) is split into N part beams by a beam splitter (2). After passing through the spectral filters (3) the remaining power is determined by broad band detectors (4). They are essentially binary filters with a 0 or 1 output depending on the wavelength. The determination of the spectral part beams transmitted is by using the algorithm described in the patent.

USE - To detect changes in substance mixtures.

ADVANTAGE - With the use of binary filters changes in the substances produce maximum changes in the signal vectors formed from the signals from the individual detectors.

DESCRIPTION OF DRAWINGS - The figure shows a block diagram of a method to the present invention.

Substance mixture 1

Beam splitter 2

Spectral filters 3

Detectors 4

Spectral power $P(\lambda)$

Dwg. 1/0

ES ESI

FA AP; GI

MC EPI: S03-E04A

L110 ANSWER 3 OF 20 WPIN (2) 1992 THOMSON DERWENT

AH 2001-027011 [78] WPIN

CF 2001-000039 [70]

DHN NL2001-467412 DNO 02001-187018

T1 Multiple filter photometer used for determining small concentration changes in a multiple component mixture comprises wide band detectors for receiving the transmitted or remitted radiation with filters.

DC J04 S03

IN EBPICH, B; MUELLER, G

PA (BECT) BECTON DICKINSON & CO; (LASE-N) LASEF & MEDICIN
 TECHNOLOGIE GMBH

CYC J1

PI DE 20018941 A1 2001101: (20017:0)* 12p G01N021-25

WO 2001079815 A1 20011021 (20017:0) EN G01N021-31 ---

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
 NL OA PT SD SE SL TC TR TS UC UY

W: AE AG AL AM AT AU AZ BA BB BG BR BY EC GA GH GI GO GR GU GZ HK DM
 DE EE EG FI GB GD GE GH GM HE HY ID IL IN IS JP KE KG KP KR KZ LC
 LK LE LI LT LU LV MA MD MG ME MI MW MX NZ NO NP PL PT RO RU SD SE
 SI SJ SK SL TH TM TR TT TD VA UG US VE VH VT ZA ZW

AU 2001065863 A 20011030 (200219) G01N021-31 ---

ALT DE 20018941 A1 DE 2000-10018941 20000417; WO 2001079815 A1 WO 2001-EP3934
 20010406; AU 2001065863 A AU 2001-05863 2001-406

FFT A1 2001065863 A Based on WO 200179815

PPAI DE 2000-10018941 20000417; DE 2000-10018940 20000417

IC ICM G01N021-31; G01N021-31

ICS G01N003-31; G01N021-31

AB DE 20018941 A UPAB: 20000321

NOVELTY - Multiple filter photometer comprises three wide band detectors (3) for receiving the transmitted or remitted radiation. The detectors have filters (4) having different spectral binary transmissions of approximately 0 and 1.

USE - Used for determining small concentration changes in a multiple component substance mixture, e.g. the change of blood glucose in a living organism.

ADVANTAGE - The concentration changes can be exactly determined.

DESCRIPTION OF DRAWING(1) - The drawing shows a schematic view of the photometer.

thermal radiator 1

filters 4

wide band detectors 5

Wdg. 1/8

ES CPl EPl

FA AB; GI

MC CPl: J04-E01; J04-C04

EPl: S03-A01A; S03-E04A; S03-E04B; S03-E04B1A

TECH UPTX: 10011211

TECHNOLOGY FOCUS - INSTRUMENTATION AND TESTING - Preferred Arrangement:

The number of detectors used corresponds to the number of relevant different parameters of the system. The wide band filters are selected so that at least one of the filters lies in a known absorption band of the goal substance. The transmitting spectral regions of the filter are selected so that the chances in the detector signals caused by substance concentrations and surroundings parameters are maximized. A thermal radiator (1) is used as a light source.

L110 ANSWER 4 OF 10 WPIM (C) 2000 THOMSON DERWENT

AI 1001-443245 [48] WPIM

DEN NL001-327863 DEN CL001-134215

TI Near infra-red spectroscopy online process comprises analyzing constituent liquids against reference spectra data bank of binary mixtures of possible solutions.

DC J04 S03 T01

IN BORN, J; FRICKEL, H; ITZEL, H

PA (MEMO) MERCK PATENT GMBH

CYC 31

FI DE 19963561 AI 19960735 (1996148)* 13p G01N021-31

WO 2001044455 AI 19960735 (1996148) DE G01N021-31 <--

FW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE TR

W: JE US

ADT DE 19963561 AI IE 1999-19963161 19991223; WO 2001044455 AI WO 2000-EP12199 19991105

PRAI IE 1999-19963561 19991223

IC ICM G01N021-31; G01N021-35

IOS G06F017-40

AB DE 19963561 A UPAB: 19960919

NOVELTY - A near infra-red spectroscopy online process analyses the constituent parts of liquid mixtures using a calibration data bank holding the reference spectra of only binary mixtures of all the possible solutions in pre-defined quantitative steps.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for an apparatus for the above process.

Preferred Features: The pre-defined steps are at irregular intervals of 1-10%. The calculation of spectra consisting of three or four components is effected by a linear combination of the data bank's two-component spectra, taking their respective proportions into account. Evaluation is effected by direct comparison of the compressed full spectral data. Both the spectra measured and the reference spectra are characterized solely by an index number used in conjunction with a data reduction module. After data compression and spectral data point scale change and respective wave count, the sum product for each spectrum is registered, and the differences between the product sums is used for rapid identification of the match between the values measured and the data bank spectra. During data compression, the respective gradient for measured and reference values is coded as a series of numbers, by adding the descending and ascending spectral data points and string of numbers.

USE - The process analyzes the constituent fluid substances in chemical liquid effluent arising from chemical batch production.

ADVANTAGE - The process takes less than 30 seconds and is external to and does not interrupt a continuing main process. The process also provides a quantitative indication of water content.

fwg.0.3

FS CPI EPI

FA AS

MC CPI: J04-B-01A

EPI: 003-E04A5B; 303-E04A5L; 303-E04A7; T01-C06B; T01-C02; T11-E01C;
T01-E02A; T01-E02B; T01-H030E; T01-T04B1; T01-J07A

L110 ANSWER 3 OF 20 WPIN 00 2002 THOMSON DEFWENT

AN 2000-448019 [59] WPIN

DN 2000-330445 DNE 02000-134904

TI Amount of fertilizer to be applied to growing grain crop, its yield and grain quality are calculated using formula.

EC 007 P11 P13 S03

IN HOSAKA, Y; MARYAMA, H; NAKAMURA, N; SATAKE, S

PA (SATA) SATAKE CORP; (SATA) SATAKE ENG CO LTD; (SATA) SATAKE SEISAKUSHO KK

CYC 5

FI CA 2290779 A1 20000309 (200039) * EN 69 A 10011-10

AU 2044890 A 20000310 (200039) A010011-00

CH 1251737 A 20000503 (200039) A010011-00

JP 2000060077 A 20001031 (200039) 13 A010011-00

KR 2000023903 A 20000415 (200107) A010011-00

ADT CA 2290779 A1 CA 1999-2290779 19990326; AU 2044890 A AU 1999-44890

19990401; CH 1251737 A CH 1999-118561 19990409; JP 2000060077 A JP

1999-154866 19990602; KR 2000023903 A KR 1999-37531 19990901

PHAI JP 1999-154866 19990602; JP 1999-154717 19990909; JP 1999-40320

19990918

IC 10M A010021-00; A010007-00

10S A010008-00; A010010-00; G01N021-31; G06F019-00

AB CA 2290779 A USAR: 20000310

NOVELTY - The amount of fertilizer to be applied to a growing grain crop is calculated using a formula including information relating the leaf blade to specific periods in the life of the crop, fertilizer application related to these periods and information about grain quality. The leaf blade information and target yield of the crop may be entered into the formula.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the following:

(1) estimating the yield of a grain crop using a formula using the same information as the novel formula;

(2) apparatus for determining the amount of fertilizer to be applied, crop yield or crop quality, having a memory storing the novel formula, a user data input, an arithmetic calculator and an output display showing the amount of fertilizer to be applied;

(3) apparatus for providing production information of grains, comprising a memory for storing a quality related formula obtained by analyzing growth information including leaf blade formation related to specific periods of the crops, fertilizer application information, and quality information of the grains after growth, an input section, an arithmetic section and a display section; and

(4) apparatus for providing production information of grains, comprising a memory storing a yield related formula obtained by analyzing growth information, and yield information of the grains after growth, an input section, an arithmetic section, and a display section.

USE - The formula can be used to calculate the amount of fertilizer to be applied to a crop, its yield, and the grain quality of the crop (claimed).

ADVANTAGE - The grain yield and quality can be estimated accurately before harvest.

fwg.2.7

FS CPI EPI GMPI

FA AB; GI; DCN
MC CPI: C05-A01B; C06-P18; C11-C07B; C12-K04E
EPI: 307-E04A5
TECH UPTX: 10010818
TECHNOLOGY FOCUS - AGRICULTURE - Preferred Features: The leaf blade information is the content of nitrogen and chlorophyll, and leaf color. These are obtained by spectral analysis. The formula may also use information on soil quality.

L110 ANSWER 1 OF 10 WPIN (C) 1001 THOMSON DEFWENT
AN 1998-611905 [53] WPIN
INN H1998-460910
TI Circuit for compensation of background absorption in an atom absorption spectrometer.
DC S03 T01 U11 V01
IN RAEFWINDEL, W; EICHARDT, E
PA (ANAL-N) ANALYTIK JENA GMBH ANALYSENMESSGERAETE
CYC 1
PI DE 19816042 A1 19891014 (19953)* 15p H01F007-06
ADT DE 19816042 A1 DE 1998-1981604. 19980409
EPAI DE 1998-19816042 19980409
IC 11M H01F007-06
103 G01J003-41; G01N021-31; G06F007-00
AB DE 19816042 A UPAB: 19991215
NOVELTY - The circuit compensates the background absorption using the magnetic field of a electromagnet for producing the Zeemann effect in an atom absorption spectrometer. The circuit has a parallel circuit of two d.c. voltage sources (E1) connected in series and two switches (S1, S2) connected in series with the magnet coil (L) in series with a current measurement device (Em) connected between their junctions.
DETAILS DESCRIPTION - An INDEPENDENT CLAIM is also included for a method of controlling an electromagnet for producing the Zeemann effect in an atom absorption spectrometer.
USE - For compensation of background absorption in an atom absorption spectrometer.
ADVANTAGE - The voltage supply enables the coil current to follow any positive or negative demand value at maximum rate.
DESCRIPTION OF DRAWING(S) - The figure shows a simplified circuit diagram.
switches S1, S2
voltage source E1
coil L
current measurement device Em
Fig. 1
FA 507
FA AB; GI
MC EPI: C05-A01B; C06-P18; C11-C07B; C12-K04E

L110 ANSWER 2 OF 10 WPIN (C) 1001 THOMSON DEFWENT
AN 1999-508391 [41] WPIN
INN H1999-378853 LDC C1999-148413
TI On-line measurement of process stream of sugar beet, sugar cane, silage, grain, fruit, vegetables, particle board or paper.
DC D12 D14 D17 P00 J04 S03
IN ATHERTON, P G; BERDING, H; BROTHERTON, G A; GRIMLEY, S C; LETHBRIDGE, P J; HAKKINTOSH, D L; STANTON, S P; STANTON, S P
PA (SUGA-N) BUREAU SUGAR EXPERIMENT STATIONS; (SUGA-N) SUGAR NORTH LTD
CYC 34
PI W) 9944193 A1 19990706 (199942)* EN 34p G01N021-31 --
RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LJ MC MW NL
OA PT SD SE SG TG TW
W: AL AM AT AU AC BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD
GE GH GM HN HU ID IL IS JP KE KG KP KR KE LC LK LR LS LT LJ LV MD

MG MK MN MW MX NO NZ PL PT PO PU SP SE SG SI SK SL TJ TM TR TT UA
UG US UZ VN YU SW

ZA 9811727 A 19990831 (199942) 11p G01N000-00
AU 9812126 A 19990713 (199941) G01N001-31 <--
BR 9814406 A 19991010 (200055) G01N001-31 <--
AU 9817034 B 19981125 (200103) G01N001-31 <--
ADT WO 9834124 A1 WO 1998-AU251 19981117; ZA 9811783 A ZA 1998-11783 19981222;
AU 9812126 A AU 1998-12126 19981117; ER 9814406 A BR 1998-14406 19981117,
WO 1998-AU251 19981117; AU 710034 B AU 1998-12126 19981117
FLT AU 9812126 A Based on WO 9834124; ER 9814406 A Based on WO 9834124; AU
717034 B Previous Publ. AU 9812126, Based on WO 9834124
PFAI AU 1997-1155 19971213
IC ICM G01N000-00; G01N021-31
ICS G01N000-00; G01N000-00; G01N021-31; G06F000-00; G06K000-00
AF WO 9834123 A EPAB: 19941014

NOVELTY - On-line measurement of a process stream reads the infrared reflectance spectrum of the stream and processes it using a reference calibration equation.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the following:

(1) A system having a scanning head with a light source and collector of reflected light, a near infrared spectrophotometer with a monochromator for resolving the reflected light into a discrete wavelength, a database storing the equation and a computer.

(2) A method of on-line measurement where the equation statistically validates the reflectance spectrum.

(3) As (a) or (b) where the process stream is sugar cane.

Preferred Features: The parameter is fiber content, juice brix, juice polarisation, commercial can sugar, quality parameters, inorganic elements or process parameters. Where the material is sugar cane it may be at any stage from prepared cane to crystalline sugar. The spectrophotometer is insulated from temperature and vibration by connecting it to the scanning head by a fiber optic cable and the head has a vibration damping mounting. The system has several spectrophotometers, one of which acts as a standard for the system. The spectrum is 400 - 2500 nanom.

USE - (all claimed) Processing sugar beet, sugar cane, silage, grain, fruit, vegetables, particle board or paper

ADVANTAGE - Infrared spectrum measurements can be made on-line.

Dwg.076

FS CFI EPI

FA AE

MC CFI: D03-R04; D06-C; F05-A040; F06-A07; C11-C
EPI: S03-E04A5

L110 ANSWER 2 OF 10 WPIN 100 00 THOMPSON DESPENT

AN 1994-132733 [21] WPIN

DNN H1994-144374 DUC G1994-011140

TI Production monitoring using a near infrared spectrometer - comprises comparing two groups of spectra using algorithm to identify changes in product quality.

EC H05 J04 S03

IN SABY, C; SABY, C A

PA (EPAB) ELF ANTAR FRANCE; (EPAB) ELF ANTAR FRANCE CA

CYC 31

PI EP 986677 A1 19980429 (199821) 11p G01N001-31 <--
E: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
FR 9754899 A1 19980423 (199823) 11p G01N001-31 <--
CA 2217108 A 19980423 (199836) G01N001-25 <--
US 6012019 A 19990919 (200008) G01N001-31 <--

ADT EP 986677 A1 EP 1997-403493 19971021; FR 9754899 A1 FR 1996-12917
19961023; CA 2217108 A CA 1997-2217108 19971022; US 6012019 A US
1997-986436 19971023

PRAI FR 1996-12917 19961023

IC ICM G01N021-25; G01N021-31; G01N021-35
 ICS G01D018-00; G01N033-28; G01N037-00;
 G06F017-00; G06F017-10

AB EP 938677 A UPAB: 19880529

Following and surveying the function of a unit fabricating a product and/or a near infrared spectrometer fed by the product, where the spectrometer delivers spectra comprising a series of absorbance values for different wavelengths, comprises: (i) recording the spectra from the near infrared spectrometer periodically in the form of numerical data; and (ii) transforming the data from each spectrum mathematically to obtain transformed spectra. In addition the following stages are used: (i) a series of working spectra are made from the obtained spectra by choosing the wavelengths in each transformed spectrum by a method of selection; (ii) a first set of 20-50 consecutive spectra are selected from the working spectra and a second set of spectra the same size is selected so they are consecutive and out of phase to the first set; and (iii) at least one quality criterion is calculated to compare the two sets of spectra and the evolution of this criterion is followed over time.

USE - Used for monitoring industrial chemicals, petrochemicals, pharmaceuticals, foodstuffs etc..

ADVANTAGE - The process is easy to use and identifies changes in product due to dysfunctions.

Int.1/3

FS CFI EPI

FA AF; GI

MC CFI: B05-K; J04-C0

EPI: C01-A02; C01-EC4A0; C01-EC4A0B; C01-E14A; C01-E14A1; C01-E14F

L110 ANSWER 9 OF 20 WP1X (C) JOHN THOMSON DECEMENT

AN 1997-297819 [27] WP1X

DNN N1997-246132

TI Instrument for optical measurement of living body - has several light modules emitting light into several positions of body through optical fibres, with beams of light transmitted to surface of body picked up at several positions by photodetectors.

DC F-1 S03 S05 T01

IN KOICUMI, H; MAKI, A; YAMASHITA, Y

PA (HITA) HITACHI LTD; (PO12-I) KOICUMI H; (KAMI-I) MAKI A; (YAMA-I) YAMASHITA Y

CYC 5

PI WO 9718755 A1 19970629 (199729) CA 1 up AC1B005-14
 W: CA DE GB US

JP 98138825 A 19970527 (199731) 13p AC1B005-14

JP 98143894 A 19970610 (199735) 11p AC1B005-14

JP 98149902 A 19970610 (199735) 11p AC1B010-00

GB 2311854 A 19971007 (199743) 13p G01N021-17 ---

DE 19681107 T 19971011 (199804) AC1B006-00

GB 2311854 B 20000322 (200115) G01N021-17 ---

US 6240309 B1 20010519 (200133) AC1B005-01

US 2001018554 A1 20010909 (200133) AC1B001-00

CA 2218703 C 20011009 (200103) EN AC1B003-14

ADT WO 9718755 A1 WO 1996-3P3365 19961115; JP 98135825 A JP 1996-299542 19951117; JP 98143894 A JP 1996-314196 19961201; JP 98149902 A JP 1996-314993 19961130; GB 2311854 A WO 1996-3P3365 19961115; GB 1997-13004 19970619; DE 19681107 T DE 1996-1961107 19961115; WO 1996-3P3365 19961115; GB 2311854 B WO 1996-3P3365 19961115; GB 1997-13004 19970619; US 6240309 B1 CIP of US 1995-539871 19961006; WO 1996-3P3365 19961115; US 1997-875381 19970429; US 2001018554 A1 CIP of US 1995-539 71 19951006, Cont of US 1997-875081 19970429; US 2001-4459 20010107; CA 2210703 C CA 1996-2210703 19961115; WO 1996-3P3365 19961115

FDT GB 2311854 A Based on WO 9718755; DE 19681107 T Based on WO 9718755; GB 2311854 B Based on WO 9718755; US 6240309 B1 CIP of US 560309, Based on WO 9718755; US 2001018554 A1 CIP of US 560309, Cont of US 6240309; CA

1010703 C Based on WO 8718755

PRAI JP 1995-14195 19951201; JP 1995-299542 19951117; JP 1995-311992 19951130

REP AU 641684; DE 4314445; DE 431-213; DE 4393335; DK 136392; DK 44693; DK 45704; EP 187539; EP 319158; EP 614645; EP 619055; FI 951673; JP 1262839; JP 5241414; JP 590512; JP 4160144; JP 5102186; JP 5111438; JP 5261110; JP 5211231; JP 61932669; JP 6181449; JP 62281815; JP 6241944; JP 63005234; JP 63805445; JP 710364; JP 7118595; JP 7119384; JP 7294168; JP 8233727; JP 819408; NO 351730; US 4576173; US 4890311; US 5408194; WO 8600514; WO 891223; WO 9401299; WO 9410901

IC 1CM A61B001-00; A61B005-05; A61B005-14; A61B006- 0; A61B010-03; G01N021-17

1CC A61B001-00; A61B005-00; A61B006-03; F01N021- 05; G01N021-27; G01N021-31; G01N033-49; G06F003-00

AE WO 8718755 A HEAB: 15372741

The optical measurement instrument has a light source (1) which includes several light modules (2(1) to 2(16)) which emit intensity-modulated beams of light at different frequencies through optical fibers (3-1 to 3-16) so that they can be introduced into the living body (9) at several points. The beams of light passing through the body are picked up on the surface of the living body (9) and guided to photodetectors (11-1 to 11-16) through optical fibers (10-1 to 10-16).

The signals from the photodetector (11-1 to 11-16) are inputted to a lock-in amplifier module (12), where the intensity of the return beam detected by each of the photodetectors and having the same modulation frequency as that of its corresponding input beam is selectively measured. The intensities of beams of light picked up at several positions are processed by a data processor (16).

ADVANTAGE - Internal information of living body for several pick-up positions can be obtained without crosstalk.

19950407

EP EPI GMEY

EA AB; GI

MC EPI: S03-E04C3; S03-D01X; T01-F 6A

L110 ANSWER 10 OF 20 WPIN (C) 2002 THOMSON DEWEENT

AN 1997-214213 [20] WPIN

ENN 1999-177184

TI Controlling working of analyser and fabrication unit for control laboratories e.g. petroleum - using multi variate calibration of master analyser, periodic standardisation of slave analysers and calibration transfers between analysers.

DC S03

IN FIDON, P; SARY, C A

FA (EPA) ELF ANTAR FRANCE; (EPA) ELF ANTAR FRANCE SA

CYC 11

FI EP 163511 AL 19970416 (199711) FR 10p G 11021-17
E: BE CH DE DK ES GR IT LI NL SE

FR 1133323 A1 19970416 (199711) G 11021-17

CA 1137943 A 19970417 (199711) FR G 11021-17

JP 0917-756 A 19970711 (199711) 10p G 11021-17

EP 168821 A 19971012 (199819) G 11021-17

IL 119427 A 20000131 (200015) G 11021-17

US 6128544 A 20001003 (200015) G06F019-00 <--

ADT EP 168522 A2 EP 1996-402187 19961015; FR 173028 A1 FR 1995-12067

19961015; CA 1187943 A CA 1996-1187943 19961015; JP 09173756 A JP

1996-304453 19961016; EP 168522 A3 EP 1996-402187 19961015; IL 119427 A IL

1996-119427 19961015; US 6128544 A US 1996-732117 19961015

PRAI FR 1995-12067 19961015

REP NG-SK.Pub; 1.Ch1.Ref; US 4366644; US 5243546; US 5459677

IC I-31 G01N019-00; G01N021-00; G01N035-00; G06F019-00

ICS G01N021-00; G01N021-31; G01N021-35; G06F015-18;

G06F017-10

ICA G01N021-27

AB EP 764532 A UPAB: 19970516

The method of following and controlling the operation of a slave analyser linked to a fabrication unit uses an operation of multivariate calibration of the master analyser and periodic standardisation operations of signals delivered by the slave analyser fed by products of standardisation.

Calibration transfer operations between the master and slave analysers are undertaken including the calculation of parameters associated with a calibration transfer algorithm and the choice of a control indicator. The evolution of the indicator is reviewed at the start of each periodic standardisation operation and used to check the correct working of the slave analyser and fabrication unit.

USE: ADVANTAGE - Also for research laboratories and manufacturing units in chemical, pharmaceutical, cosmetic, food and agricultural industries. Method enables causes of drift and malfunction to be identified.

Dwg. 0-0

FC FBI

FA AB

MC FBI: 001-E04A1; 301-E04T

L110 ANSWER 11 OF 20 WPIX (C) 1992 THOMSON DEWENT

AI 1994-351270 [44] WPIX

DIN I1994-275640

T1 Instantaneous spectroscopic analysis system for fluid - uses correlation of spectrum produced with those held in memory to determine relative concentration of various products in fluid sample.

DC S03 T01

IN PACHINGER, C; MARTIN-BOUYER, N; NAFFRECHOUX, E; SUPIL, J

FA (UYSA-10) UNIV SAVOIE

CYC 1

FI FF 2764650 AI 19941104 (199444)* 16p G01N021-27

WO 9425837 AI 19941106 (199444) EP 16p G01F013-28

EP 647316 AI 19950412 (199519) EP G01F013-28

US 5528363 A 19960618 (199630) 6p G01F013-18

ALT FF 2764650 AI EP 1993-5104 19930427; WO 9425837 AI WO 1994-FR478 19940427; EP 647316 AI EP 1994-914451 19940427; WO 1994-FR478 19940427; US 5528363 A WO 1994-FR478 19940427; US 1994-360744 19941222

FDT EP 647316 AI Based on WO 9425837; US 5528363 A Based on WO 9425837

FEAI EP 1993-5104 19930427

REP 06Jnl.Ref; EP 368500; EP 369057; EP 510122; GB 2217006; WO 9013810

IC ICM G01C003-18; G01C003-28; G01N021-27

ICS G01C003-50; G01N021-31; G06F015-20

AB FF 2764650 A UPAB: 19941223

The detection and identification system operates by illumination of the medium being examined by a monochromatic or polychromatic light wave (A), and processing of the resulting emission, absorption or reflection spectrum. This is achieved with a plane field monochromator (5) and a transformation circuit (6) producing analogue or digital signals.

The circuit is coupled to a central processing unit (7) analysing the spectrum produced for comparison and correlation with spectra stored in memory. As a result of this comparison, the concentration of various physico-chemical products within the medium may be determined.

ADVANTAGE - System provides instantaneous analysis of composition of fluid.

Dwg. 1/1

FS FBI

FA AB; G1

MC FBI: 001-E04A1; T01-J06A

ABEQ US 5528363 A UPAB: 19960731

A compact portable device capable of operating in a hostile environment to carry out qualitative and quantitative identification of one or of a plurality of physicochemical entities contained in a sample capable of

producing a spectra under excitation by electromagnetic waves that includes:

a portable compact casing that shields equipment housed therein from magnetic fields, electric fields, and external pressure variations;

a polychromator having an optical path within said casing to which emissions, absorption or reflective spectrum of a liquid, solid or gas sample are transmitted and analysed, having said spectrum being decomposed into a sequence of signals having discrete variation in wavelength;

detecting means for detecting said discrete wavelength signals positioned on the optical path of the polychromator;

conversion circuit means coupled to said detecting means for converting said discrete wavelength signals into electrical signals; and,

processing means having a plurality of standard spectra represented of known entities in memory for analyzing the spectra embodied in said electric signals and to decorrelate and compare the spectra with that stored in memory and to determine the nature and concentration in said sample.

Dwg.171

L110 ANSWER 12 OF 30 WPIX CC, 2002 THOMSON DERWENT

AN 1994-279897 [14] WPIX

DNN N1994-110459 RUC C1994-12779.

TI Optical measuring unit for modifications in reactive substance in transparent cell - passes light through cell and then through filters to brightness measuring elements.

DC J04 S03

IN DELIGNIERES, R; DUEAND, C

FA (INSE) INST FRANCAIS DU PETROLE

CYC 19

EI WO 9418543 A1 19940516 (199404) 18p G01N011-31 8--

FW: AT BE CH DE DK EC FR GB GR IE IT LU MC NL PT SE

W: CA JP US

FR 2731316 A1 19940612 (199404) 18p G01N011-31 8--

EP 685127 A1 19940127 (199304) 10p G01N011-31 8--

R: BE DE GR IT NL SE

JP 06501394 W 19940217 (199304) 18p G01N011-79

US 5680220 A 19971011 (199704) 10p G01N011-31

EP 685127 B1 19960216 (199404) 10p G01N011-31 8--

R: BE DE GR IT NL SE

DE 6941331 E 19981021 (199404) G01N011-31 8--

AUT WO 9418543 A1 WO 1994-FR120 19940131; FR 2701316 A1 FR 1043-1513 19930209;

EP 685127 A1 EP 1994-905765 19940131; WO 1994-FR120 19940131; JP 06501394

W OF 1994-51771 19940131; WO 1994-FR120 19940131; US 5680220 A WO

1994-FR120 19940131; US 1995-30773 19970127; EP 685127 B1 EP 1994-905765

19940131; WO 1994-FR120 19940131; DE 6941331 E OF 1994-51771 19940131;

EP 1994-905765 19940131; WO 1994-FR120 19940131

FMT EP 685127 A1 Based on WO 9418543; JP 06501394 W Based on WO 9418543; US

5680220 A Based on WO 9418543; EP 685127 B1 Based on WO 9418543; DE

6941331 E Based on EP 685127, Based on WO 9418543

FRAT FR 1993-1513 19930309

REP US 5680220; US 3615342

IC ICM G01N021-35; G01N021-31; G01N011-79

ICG G01N003-46; G01N003-37; G06F015-46

AB WO 9418543 A UFAB: 19941013

The unit consists of at least one light source (1) with two optical branches (11,12) which allow the incident light to be passed selectively through the cell (3), an optical system (7,8) to direct the emergent light rays through selective optical filters (F1, F2, F3) of different wavelengths, and measuring elements (D1, D2, D3) for the intensity of light passing through the filters.

The emergent light rays from the cell and a neutral filter (5) can be passed selectively through the filters by means of two shutters (01,02) and a switching system (M,I). The unit also incorporates a control

assembly (9) with a command processor (10), a signal acquisition unit (11), and an interface assembly (12).

USE/ADVANTAGE - Appts. can be used for determining the pH of a substance. More precise and reliable results are obt'd.

lwp.1/6

FS CFI EPI

FA AB; GI

MC CFI: J04-C

EMI: J04-E04A5

ABEQ US 5060210 A UPAB: 19971210

A device for optically measuring modifications in a reacting substance contained in a transparent cell, comprising

a single light source provided with an electric supply voltage and a specified light spectrum, a first optical circuit, a first optical shutter arranged in said first optical circuit, a second optical circuit, a second optical shutter arranged in said second optical circuit, an optical diverter in said optical circuits for diverting incident light from the single light source through the cell and through a reference medium to an optical node, an optical separator for directing light from the optical node to three other optical circuits, a set of three selective filters arranged respectively in the three other optical circuits, a first selective filter from the set being centred on a first wavelength corresponding to an isobestic point of the reacting substance, a second selective filter from the set being centred on a wavelength in a part of the light spectrum where the reacting substance is the most sensitive and a third selective filter from the set being centred on another part of the light spectrum where the reacting substance is the least sensitive, a measuring means for respectively measuring the light emanating from the three other optical circuits, including a set of three detectors for respectively detecting light passed by each of the set of three filters and producing output signals representing the detected light, an electric power supply for providing the electrical supply voltage, a controller and an electric switching means controlled by the controller for connecting intermittently the three detectors to the controller, for connecting the single light source to the power supply, and for selectively switching the optical shutters.

lwp.1/6

L110 ANSWER 13 OF 30 WPBX (C) 1992 THOMSON DERWENT

AN 1994-101101 [15] WPBX

DUN N1994-15-047

TI Spectrometer with dynamically coded components - has data carriers, readers, writers and central computer for coded data characterising replaceable components.

DC S03

IN KAPP, H; SIMON, A; WEIL, J

PA BRUK-IG BRUKER ANALYTISCHE MESSTECHNIK; (BRUK-N) BRUKER ANALYTISCHE MESSTECHNIK GMBH

CYC

PI DE 4241905 A1 19940616 (199405) 8p G01J003-02

DE 4241905 C2 19950116 (199505) 7p G01J003-02

US 5557544 A 19960917 (199643) 7p H01J003-02

ADT DE 4241905 A1 DE 1993-4241905 19921211; DE 4241905 C2 DE 1993-4241905 19921211; US 5557544 A US 1993-164303 19931209

PRAI DE 1992-4241905 19931211

IC ICM G01J003-02; H01J003-02

ICS G01N021-31; G01N024-08; G01N024-10; G01N027-62; G02B027-00; G06F013-00

AB DE 4241905 A UPAB: 19940616

The spectrometer has a central computer, a radiation source, detector, beam divider, filter and external measurement probe. The replaceable components each have a readable data carrier (7) with coded data contg. the parameters characterizing each component. The data carrier is a chip,

esp. an EPROM or a flash-ROM.

The data carrier can be written into and contains variable, time dependent data concerning the prewritten and current characteristics of each replaceable component, e.g. operating duration, wear parameters or calibration curves. The central computer decodes the data and derives deviations.

USE/ADVANTAGE - Exp. for infrared analytic spectrometer, but also NMR, ESR, ICR, or mass spectrometer. Dynamically coded components can be used with great flexibility at other points in same spectrometer or in other spectrometers.

Dwg.17/

FS EPI
FA AB; GI
MC EPI: S03-A02B; S03-B04A; S03-B07; S03-E11A
ABEQ DE 424105 C UFAB: 12950301

An analytical, esp. IR, spectrometer has a CPU (3), fixed components and replaceable components (5) including a readable data carrier (7) with coded data on component parameters. These data can be read and transmitted (8a) to the CPU which has a decoding and decision-making programme.

One or more devices input variable time-dependent data into the data carrier on the actual condition of the replaceable components. The CPU has a programme to control the data writing device and to automatically match the data on changed component parameters on its data carrier.

ADVANTAGE - Greater component flexibility at different positions in the same or other spectrometers.

Dwg.17/

ABEQ US 5557544 A UFAB: 12961025
An FTIR spectrometer comprising:

an interchangeable optical component;

readable data medium means integral with the interchangeable optical component and adapted for storing encoded data concerning at least one of a history and a changeable current property of the interchangeable optical component and adapted for storing non-changeable encoded data;

read/write means connected to the data medium means for reading encoded data from and for writing encoded data to the data medium means;

a central computer adapted to decode the encoded data and to process decisions on the basis of decoded data and adapted to process the encoded data for controlling the read/write means to automatically update changed parameters of the interchangeable optical component;

interface means connected between the read/write means and the central computer for transferring the encoded data to and from the central computer; and

sensor means, communicating with the read/write means, for detecting changes in the encoded data and for generating data signals in response to the detected changes in the encoded data.

Dwg.17/

L110 ANSWER 14 OF 20 WPIX (C) 2002 THOMSON DERWENT

AN 1993-218893 [32] WPIX

DUN 11993-199136

TI Instrument for non-destructive measurement of material properties - uses data obt'd. from material by sections of electromagnetic spectrum to determine material properties by data-fusion analysis.

DC S03 T01 K11

IN ESCTERGAR, E P

PA (SGII-H) SGI INT

CYC 18

PI WO 9315470 A1 19930805 (199312)* EN 32p G06F015-46 ---

RW: AT BE CH DE DK ES FR GB GR IE IT LU MC NL OA PT SE

W: AU BR CA JP KE NZ PL RO RU UA

AU 9335940 A 19930901 (199310) G06F015-46 ---

US 5291422 A 19940301 (199409) 13p G01N021-47

JP 07507133 W 19950803 (199539) 10p G01N022-00

ADT WO 9315470 A1 WO 1993-US704 19930126; AU 9355940 A AU 1993-35940 19930126;
US 5291411 A US 1993-826720 19930128; JP 07507133 W JP 1993-513395
19930116, WO 1993-US704 19930126

FET AU 9355940 A Based on WO 9315470; JP 07507133 W Based on WO 9315470

FFAI US 1992-8.6780 19930128

REP US 5475720; US 4616817; US 4616819; US 5127262

IC ICM G01N021-47; G01N021-08; G06F015-46

ICS G01N021-31; G01N021-21

AE WO 9315470 A UPAB: 19940510

Samples (14) are carried by conveyor belt (16) past an array of sensors (20), each of which stimulates electromagnetic spectrum from the samples and detects that radiation. Each sensor has an associated microprocessor controller (22) which feeds detected data to a data processing computer (18). The data processor merges the data from the sensors by matrix techniques and passes the results to the user via a communication path (16). The user can also supply instructions and additional data e.g. from other forms of measurement to the data processor.

USE/ADVANTAGE - Analysis of e.g. coal, food products, wood, cement, petroleum. Multiple measurements performed automatically on a single sample eliminate need for multiple samples, increase accuracy and save both time and effort. Properties difficult to measure directly may be measured indirectly more easily and the required result obtd. by data fusion techniques.

Dwg.1/4

Dwg.1/4

FS EP1

FA AB; GI

MC EP1: S03-E04A; S03-E05; S03-E06; S03-E07; S03-E14E1; T01-C08A; X11-A09

AEQ US 5291411 A UPAB: 19940418

The non-destructive, non-contacting instrument detects electromagnetic radiation from the materials across a wide range of the electromagnetic spectrum, and combines these diverse data to derive the material property values desired.

In particular, the properties detectable through particle magnetic resonance, spectroscopy of light in the infrared-visible-ultraviolet range, and detection of X-ray and gamma ray radiation may be included in the instrument. Sensors detect each wavelength band of electromagnetic radiation, and data from all these sensors are merged in a central data processor to evaluate the material properties of interest.

USE/ADVANTAGE - Provides material property measurements quickly and automatically, using single sample of test material. Incorporates data fusion to enable information about material to be derived by correlation of disparate sensor data, with minimal human intervention required.

Dwg.1/4

L110 ANSWER 15 OF 30 WP1X 09 1002 THOMSON DERWENT

AN 149C-116107 [15] WP1X

TI Spectroscopic determin. of one constituent in fluid mixt. - such as oil content of wax, suitable for on-line or batch measurement, avoiding need for dilution.

DC H02 001 004 S03

IN CRIMENTI, R J L; HALPERN, G M

PA (ES:00) EXXON RES & ENG CO

CYC 6

PI EP 479472 A 14420408 (199311)* 1p

R: DE FR GB IT

CA 2059108 A 19920317 (199303) G01N021-35

EP 479472 A3 19930510 (199301)

IT 5361115 A 19940401 (199413) 1p G06F015-20 <--

EP 479472 B1 19950614 (199513) EN 4p G01N021-25

R: DE FR GB IT

DE 69110300 E 19950322 (199504) G01N021-35

CA 2059108 C 20010413 (200157) EN G01N021-25

ADT CA 2050108 A CA 1991-2050108 19910828; EP 479472 A3 EP 1991-308682
19910924; US 5301125 A US 1990-588643 19900926; EP 479472 B1 EP
1991-308682 19910924; DE 69110390 E DE 1991-610390 19910924, EP
1991-308682 19910924; CA 2050108 C CA 1991-2050108 19910828

FDT DE 69110390 E Based on EP 479472

PEAI US 1990-588643 19900926

REP No-SR.Pub; DE 3615490; GB 2020009; US 4449819; EP 3623499

IC ICM G01N011-25; **G06F015-20**

ICS G01N011-27; **G01N021-31**

AE EP 479472 A UPAB: 19931118

Process is for spectroscopic determination of amt. of one constituent of a fluid mixt. in another constituent or in the fluid mixt. itself, following sepr. of the mixt. into its constituents, in which a spectroscopic determination of the amount α_N of the Nth constituent of a fluid mixt. 0 in another constituent n of the mixt. following the sepr. of the mixt. into M constituents 1, ...M (where N is sup to M) and where, due to imperfect sepr., the amt. α_N of constituent N remains present with separated constituent n, comprises (a) determining the absorptivity a_N of constituent N at a selected wavelength, or at multiple wavelength, across a selected wavelength range, in which constituent N exhibits light absorption; (b) determining the absorptivity a_n of the other constituent n with the amt. α_N of constituent N present at the same selected wavelength or by the same multiple wavelengths; and (c) determining the amount α_n of the one constituent N present with constituent n from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities a_n and a_N where the absorptivities are expressed solely as the ratio of a_n/a_N .

USE/ADVANTAGE - The method is suitable for measuring the oil content of wax, and is suitable for on-line measurement or batch measurement, as circumstances require, the method avoiding the need for dilution. The use of the method for determining the oil content of wax obtd. by dewaxing of oil boiling in the lubricating oil range is specifically claimed.

5/6

ress

FS CFI EPI

FA AE; GI

MC CFI: N05-K; J04-B01A

EPI: S03-B04A5; S01-B04B1A

ABEQ EP 479472 A UPAB: 19931006

Process is for spectroscopic determination of amt. of one constituent of a fluid mixt. in another constituent or in the fluid mixt. itself, following sepr. of the mixt. into its constituents, in which a spectroscopic determination of the amount α_N of the Nth constituent of a fluid mixt. 0 in another constituent n of the mixt. following the sepr. of the mixt. into M constituents 1, ...M (where N is sup to M) and where, due to imperfect sepr., the amt. α_N of constituent N remains present with separated constituent n, comprises (a) determining the absorptivity a_N of constituent N at a selected wavelength, or at multiple wavelength, across a selected wavelength range, in which constituent N exhibits light absorption; (b) determining the absorptivity a_n of the other constituent n with the amt. α_N of constituent N present at the same selected wavelength or by the same multiple wavelengths; and (c) determining the amount α_n of the one constituent N present with constituent n from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities a_n and a_N where the absorptivities are expressed solely as the ratio of a_n/a_N .

USE/ADVANTAGE - The method is suitable for measuring the oil content of wax, and is suitable for on-line measurement or batch measurement, as circumstances require, the method avoiding the need for dilution. The use of the method for determining the oil content of wax obtd. by dewaxing of oil boiling in the lubricating oil range is specifically claimed.

5/6

ABEQ US 5301125 A UPAB: 19940517

When separating a fluid mixture into fractions, the amount of the Nth constituent remaining in another constituent n is determined by measuring the light absorption of N and n and determining the amount of N in n using an expression in which the absorptions are expressed solely as the ratio a_N/a_n . The method is partic. for determination of the entrained oil content of wax resulting from separation of a waxy raffinate into dewaxed hydrocarbon oil boiling in the lubricating oil range, with solvent added to samples of wax and oil fractions before the determination.

ADVANTAGE - Simplifies determ. and can be used on-line or in a batch process.

Dwg. 378

ABEQ EP 474472 B (FAB: 19960721)

A method for the spectroscopic determination of the amount a_N of the Nth constituent of a fluid mixture n in another constituent n of the mixture following the separation of said mixture into N constituents 1,...,M (where n, N is less than M) and where, due to imperfect separation, said amount a_N of constituent N remains present with separated constituent n, said method comprising the steps of:- (i) determining the absorptivity a_N of separated constituent N at a selected wavelength, or at multiple wavelengths, across a selected wavelength range, in which constituent N exhibits light absorption; (ii) determining the absorptivity a_n of said another separated constituent n with said amount a_N of constituent N present at the same selected wavelength or at the same multiple wavelengths; and (iii) determining the amount a_N of said one constituent N present with constituent n from a mathematical expression which contains, or is equivalent to an expression which contains, the absorptivities a_n and a_N where the absorptivities are expressed solely as the ratio a_N/a_n .

Dwg. 378

L110 ANSWER 16 OF 20 WFIX OF 1991 THOMSON DERWENT

AN 1991-311,06148 WFIX

DNN 11991-139309

TI Atomic absorption spectrophotometer, calibration method - uses comparative technique for determining constituent of sample by measuring and storing several standards nominal values.

DC 381 S03 T61

IN MASSART, D; MASSART, E J L

FA (PHIG) PHILIPS GLOEILAMFENFAB NV; (PHIG) PHILIPS ELTEN & ASSOC IND LTD; (PHIG) KONINK PHILIPS ELECTRONICS NV; (PHIG) PHILIPS ELECTRONICS UK LTD; (PHIG) US PHILIPS COFF

CYC #

FI EP 453036 A 19911013 (199144)*

F: CH LE FE GB LI

GB 214111 A 19911013 (199144)

AU 6175073 A 19911013 (199150)

US 5210778 A 19930511 (199310)

AU 648545 E 19940418 (199411)

US 5552997 A 19960308 (199641)

EP 453036 F1 19991201 (200001) EN

F: CH DE FE GB LI

DE 69131906 E 20000105 (200009)

JP 0421270 E 20000818 (200044)

EP 453036 A EP 1991-100380 19910415; GB 2245211 A GB 1990-8313 19900410;

US 5210778 A US 1991-685106 19910412; AU 648545 E AU 1991-75073 19910418;

US 5552997 A Cont of US 1991-685260 19910412, Cont of US 1991-976634

19911116; US 1994-175000 19940714; EP 453036 B1 EP 1991-100380 19910415;

DE 69131906 E DE 1991-61806 19910415; EP 1991-209380 19910415; JP 0421270

B2 JP 1991-115339 19910418

FDT AU 648545 B Previous Publ. AU 6175073; US 5552997 A Cont of US 5210778; DE 69131906 E Based on EP 453036; JP 0421270 B2 Previous Publ. JP 04210834

FFAI GB 1990-8312 19900410

REP 4.Jnl.Ref; A...9150; LE 3406123; NoSE.Pub

IC G01N021-31; G01N023-06; G06F015-20
 ICM G01N018-00; G01N021-27; G01N021-31; G12B012-00
 ICS G01N023-06; G06F015-20; G06F019-00
 ICA G01N023-20; G01N023-22
 AB EF 454036 A UPAB: 10930928

The method of calibrating an analytical instrument which uses a comparative technique for determining a constituent of a sample involves measuring a characteristic of several standards having different nominal values. The measured characteristic is stored in association with its corresponding nominal values. A best straight line is determined using statistical techniques on the stored values. The quality of the calibration line is determined. When the quality of the calibration line is not acceptable, the slope of the line joining each of the stored measured characteristics and nominal values to the origin is determined. It is also determined whether the slopes have a given order.

The method comprises determining the slope of each stored measured characteristic and nominal value with respect to the first measured characteristic and nominal value. The curvature of the calibration line is determined if the slopes have a given order. If the slopes have not a given order it is indicated that the calibration line does not pass through the origin.

ADVANTAGE - Enables determination of possible causes of lack of quality of calibration line, by finding whether or not slopes are random so determining whether the problem is in precision of measurement of standards or quality of standards, or whether the points do not represent straight line through origin, but represent curved line.

1/7

FS EPI
 FA AB; G1
 MC EPI: S02-K09; S03-E04D; T01-J
 ABEQ US 5310778 A UPAB: 10931113

The atomic absorption spectrophotometer has advice for measuring the absorbance of a number of standards of known concentration (100) and plots the measured absorbance against concentration (101). A straight line is fitted to the plotted points (102) and a quality coefft. calculated (103). If the quality coefft. is acceptable (104) the calibration line is used for measurement of samples (105). If not, then the slope of the line joining each point to the origin is determined and if the slopes are random (107) then a robust regression technique is used to fit the calibration line (106). If outliers are then detected (109) it is determined which points are outliers (110) and appropriate action taken, for example to restrict the range if the last point(s) is/are outliers (111).

2/7

If the slopes determined in step (106) are not random, then, provided more than four points remain (112), the slope of each point with respect to the first point is determined (114). If they are again not random (115), then a curved calibration line is diagnosed while if they are random, a straight line not passing through the origin is diagnosed (117). In atomic absorption spectroscopy a straight line not passing through the origin indicates a problem with the blank solution, for example, contamination.

USE - For determining chemical properties of sample.

Dwg. 2/7

ABEQ US 5552997 A UPAB: 10961011
 A method of calibrating an X-ray spectrometer comprising the steps of
 (a) generating x-rays from at least one standard sample,
 (b) measuring intensities of said x-rays for different concentrations of said standard sample-,
 (c) forming a representation of intensity versus concentration for each measured value of intensity with said different concentrations,
 (d) determining if a best straight calibration line can be formed from said representation, and if not

(c) changing instrument parameters and/or sample preparation to achieve said best straight calibration line, and

(d) repeating said steps (a)-(c) to calibrate said x-ray spectrometer.

Dwg. 1/7

L110 ANSWER 17 OF 20 WPIX (C) 2001 THOMSON DEFWENT

AN 1983-001453 [01] WPIX

DIN N1989-001581

TI Optical interference, absorption or scatter determining appts. - has measured values of electromagnetic radiation intensity correlated with different sets of known values derived from model.

DC S02 S03 T01

IN EDGAR, R F

PA (INFR-N) INFRARED ENG LTD; (INFR) INFRARED IND INC

CYC 6

PI GB 2206429 A 19830105 (198301) 13p

EP 299646 A 19830118 (198303) EN

E: BE DE FR GB IT

US 4952061 A 19900828 (199037)

EP 299646 B1 19930505 (199318) EN 16p G01N021-31 ---

E: BE DE FR GB IT

DE 3880748 G 19930609 (199324) G01N021-31 ---

ALT GB 2206429 A GB 1987-15608 19870702; EP 299646 A EP 1988-305915 19880629;

US 4952061 A US 1988-211708 19880627; EP 299646 B1 EP 1988-305916

19880629; DE 3880748 G DE 1988-3880748 19880629; EP 1988-305915 19880629

FMT DE 3880748 G Based on EP 299646

PRAI GB 1987-15608 19870702

REP A: 1983; DE 2426598; EP 230305; No-GR.fab; SU 1234508; US 4555767; 91Jnl.Ref

IC G01B011-02; G01N021-31; G06F000-01

ICM G01N021-31

ICS G01B011-02; G01N021-35; G01N021-41; G01N021-47; G01N021-38;

G06F000-01

AE GB 2206429 A UPAB: 19830923

Electromagnetic radiation is transmitted through or reflected from a sample. The radiation includes at least two spectrally different components so that at least one of the components is subjected to optical interference, absorption or scatter. The components are transmitted through or reflected from the sample by respectively different amounts.

The transmittance or reflectance of the sample for each of the components is measured to derive respective measured values, before correlating by either a zero dependent correlation function (Z), or a residual function (Nres). Known values having an optimum correlation with the measured values are selected, with the selected values representing the property, or the identity of the sample which is sensed or to be determined.

ADVANTAGE - Both functions are unaffected by gain factors, thus avoiding any need to determine and to maintain absolute sensitivities of optical detectors, provide greater variation of correlation and increasing precision with which optimum correlation can be determined, especially when either the measured values, or known values are subject to error.

Q/3

FS EPI

FA AB; GI

MC EPI: S02-A03A; S03-E04; T01-S04B

ABEQ EP 299646 B UPAB: 19831112

A method of sensing or determining one or more properties or the identity of a sample in which electromagnetic radiation is subject to optical interference, absorption or scatter, the method comprising the steps of: (a) causing electromagnetic radiation to be transmitted through, or reflected from said sample, said radiation including at least two spectrally different components so that at least one of said components is

subjected to said optical interference, absorption or scatter and so that said components are transmitted through, or reflected from said sample by respectively different amounts; (b) measuring the transmittance or reflectance of said sample for each of said components to derive respective measured values; (c) correlating by means of either a zero dependent correlation function 'S', or a residual function 'Nres', respectively defined by: said measured values of transmittance or reflectance with different known values representing or relating to either different values of a property of a known material, or different values which are characteristic of different known materials; and (d) selecting the known values having an optimum correlation with said measured values, the selected known values representing the property, or the identity of the sample which is sensed or to be determined.

1/3

Dwg. 1/3

ABEQ US 4952061 A UPAB: 199309L

Sets of measured values of the intensity of electromagnetic radiation, which has been subject to optical interference, absorption or scatter by a sample, are correlated with different sets of known values derived from either model of the optical properties of the sample, or from an analogue technique, correlation is by means of either a zero dependent correlation function, or a normalized residual function.

Both functions are unaffected by gain factors, avoiding any need to determine and to maintain absolute sensitivities of optical detectors, provide greater variation of correlation than with techniques employing a conventional correlation coefficient.

ADVANTAGE - Increased precision with which the optimum correlation can be determined, when either the measured value, or known values are subject to error. Reduced computing time.

L110 ANSWER 18 OF 38 WPIX (C) 2003 THOMSON DEWENT

AN 1988-310508 [41] WPIX

CR 1988-310511 [45]; 1988-310515 [41]

DNN M1988-239993

TI Examination appts. for measuring oxygenation of blood - monitors variations in fitting position of illumination side fixture by detecting light reflected from object.

DC P81 S03 S05 V87

HU HAKAMATA, H; OCAKI, T; SUZUKI, S; YAGI, S

PA (HAMM) HAMAMATSU PHOTONICS KK

CYC 3

PI EP 290772 A 19881109 (198843) EN 11p

F: DE GB

US 4901338 A 19900213 (199013) 11p

EP 190171 D1 19880714 (198813) EN 11p A61B005-00

F: DE GB

EP 190171 B1 19890714 (198813) EN 1 p A61B005-00

F: DE GB

EP 190171 B1 19890714 (198813) EN 1 p A61B005-00

F: DE GB

DE 3882172 G 19880819 (198834) A61B005-00

DE 3882174 G 19880819 (198834) A61B005-00

AOT EP 190171 A EP 1988-304130 19880506; US 4901338 A US 1988-118512 19880502;

EP 190171 B1 EP 1988-304130 19880506; EP 190171 B1 EP 1988-304130

19880506; EP 190171 B1 EP 1988-304130 19880506; DE 3882172 G DE

1988-3882172 19880506; EP 1988-304130 19880506; DE 3882174 G DE

1988-3882174 19880506; EP 1988-304130 19880506

FOT DE 3882172 G Based on EP 290.72; DE 3882174 G Based on EP 290.75

PRAI JP 1987-110461 19870508; JP 1987-675841 19870508; JP 1987-110465

19870508; JP 1987-110471 19870508

REP GB 2061496; GB 2075668; GB 2151827; GB 2593767; US 5936192; EP 123548; EP

160768; FR 2539613; GB 2064444; US 4281645

IC A61B005-00; G01N021-31; G06F015-42

ICM A61B005-00
ICS G01N021-25; G01N021-31; G06F015-42

AB EP 290272 A UPAB: 19931116

The examination appts. measures the oxygenation using near IR light of different wavelengths. The appts. has a light source controller, an illumination side fixture, a detection side fixture a transmitted light detector and a computer to control the appts. and analyse the results. The body's heartbeat period is divided into several cycles and the transmission quantities of radiation transmitted through the head or organ are accumulated at every wavelength and for every cycle.

The computer judges whether a fitting position of the illumination side fixture has been changed on the basis of the reflection light data and the output light data.

USE - Diagnosis of cerebral tissue damage.

Dwg.0/6

FS EPI GMPI

FA AB

MC EPI: 305-E04; 305-001

ABEQ TM 4401258 A UPAB: 19930923

The examination device comprises light source for sequentially emitting electro-magnetic waves with different wavelengths, with an illumination-side fixture for making the electro-magnetic waves introduced from the light source incident on a measuring object and detects reflected electro-magnetic waves from the measuring objects. A reflection light detector detects the reflected electro-magnetic waves introduced from the illumination-side fixture and outputting reflection light data. An output light detector detects emitted electro-magnetic waves from the light source and outputting output light data.

A computer system receives the reflection light data from the reflection light detector and output light data from the output light detector and judges whether a fitting position of the illumination-side fixture has been changed on the basis of the reflection light data and the output light data.

USE - For measuring the oxygenation in an object with electromagnetic wave transmission spectrophotometry.

ABEQ EP 290272 B UPAB: 19931116

An examination apparatus (1) for measuring the oxygenation in an object (40) with electromagnetic wave transmission spectrophotometry, comprising; light source means (L01-L04) for sequentially emitting electromagnetic radiation of different wavelength; an illumination-side fixture (51) for contacting the electromagnetic radiation generated by the light source means (L01-L04) with an object (40); a receiving-side fixture (52) for receiving electromagnetic radiation transmitted through the object (40); a transmitted light detection device (54) for detecting electromagnetic radiation received by the receiving-side fixture (52); and a computer system (6) for controlling the light source means (L01-L04) and the transmitted light detecting device (54) and for analysing the output of the transmitted light detecting device (54) to determine the oxygenation of the object (40). characterised in that the illumination-side fixture (51) is arranged to receive electromagnetic waves reflected from the object (40); in that the examination apparatus also includes; reflected light detection means (4) for detecting the reflected electromagnetic radiation introduced from the illumination-side fixture (51) and outputting reflected light data; and output light detection means (13) for detecting electromagnetic radiation emitted by the light source means (L01-L04) and outputting output light data; and in that the computer system (6) receives the reflection light data from the reflection light detection means (4) and the output light data from the output light detection means (13), and determines whether a fitting position of the illumination-side fixture (52) changes on the basis of the reflection light data and the output light data.

Dwg.1/6

ABEQ EP 290275 B UPAB: 19931116

An examination apparatus for measuring the oxygenation of an object by electromagnetic radiation transmission spectrophotometry, comprising: light source means (LD1-LD4, 38) for sequentially emitting electromagnetic radiation at a number of different wavelengths; an illumination-side fixture (32) for applying the electromagnetic radiation emitted by the light source means (LD1-LD4) to an object (40); transmitted light detection means (54) for detecting electromagnetic radiation transmitted through the object (40) and outputting transmission light data; a detection-side fixture (34) for receiving electromagnetic radiation transmitted through the object (40) and coupling it to the transmitted light detection means (54); and, a computer system (56) for controlling the light source means (LD1-LD4, 38) and the transmitted light detection means (54), receiving the transmission light data, and calculating the oxygenation in the object (40); characterised in that the illumination-side fixture (32) is equipped with a first indication means (33) for indicating if electromagnetic radiation is being emitted from the light source means (LD1-LD4); and the detection-side fixture (34) is equipped with a second indication means (35) for indicating if the transmitted light detection means (54) is in its operating condition; and, shapes and/or colours of the illumination-side fixture (32) and the detection-side fixture (34) are different to one another.

Dwg.12/11

L110 ANSWER 19 OF LC WPIX (C) 2001 THOMSON DEWENT

AN 1487-137535 [05] WPIX

DNN N1987-103088

TI Single optical fibre transducer driving and measuring circuit - has bi-directional couplers recording signal intensities and transmitting pulsing energy to wavelength multiplexer-demultiplexer.

DC FBI S03 S05 V07

IN MRAJCH, S R

PA (BECT) BECTON DICKINSON CO; (DECE-W) DESEFET MEDICAL

INC

CYC 14

FI EP 222555 A 19870513 (198805) EN 3p

R: BE CH DE ES FR GB IT LI NL SE

US 4936679 A 19900816 (1988023)

CA 1352131 C 19910402 (1990113)

EP 222555 B 19910813 (1988023)

R: BE CH DE FR GB IT LI LU NL

DE 3681117 G 19911002 (1990141)

ES 1026132 T3 19910415 (199006)

AU 6085077 A 19870514 (198804)

G01N021-31 <--

ADT EP 222555 A EP 1986-308430 19861029; US 4936679 A US 1985-797299 19851112;

EP 222555 T3 EP 1986-308430 19861029

FDT EP 222555 T3 Based on EP 222555

PRAI US 1985-797299 19851111

REP A61B813; No-SR.Pub; US 347444; US 3036911; US 4114604; US 4444498

IC ICM G01N021-31

ICS A61B005-00; G01N033-41

AB EP 222555 A UPAB: 19860921

The system has an energy source connected to a power supply for emitting bursts of energy at a predetermined colour frequency. Another energy source is connected to the power supply for emitting bursts of energy at another predetermined colour frequency. A wavelength division multiplexer/demultiplexer is associated with the two energy sources for receipt of the bursts of energy, and for combination and further transmission.

The discrete colour frequencies are maintained and the reflective energies are separated into individual channels of each frequency upon returns of the bursts of energy. A fibre optic device and delay unit are arranged for receipt of the combined bursts of energy.

USE/ADVANTAGE - For in vivo measurement of blood physiological

parameters. Enhances strong returns signal.

1/2

FS EP1 3MPI

FA AB

MC EPI: 303-E04A9; S05-C01; S05-D01X; V07-K04; V07-N

AREQ EP 22355 B UPAB: 19930922

The system has an energy source connected to a power supply for emitting bursts of energy at a predetermined colour frequency. Another energy source is connected to the power supply for emitting bursts of energy at another predetermined colour frequency. A wavelength division multiplexer/demultiplexer is associated with the two energy sources for receipt of the bursts of energy, and for combination and further transmission.

The discrete colour frequencies are maintained and the reflective energies are separated into individual channels of each frequency upon returns of the bursts of energy. A fibre optic device and delay unit are arranged for receipt of the combined bursts of energy.

USE/ADVANTAGE - For in vivo measurement of blood physiological parameters. Enhances strong returns signal.

1/2

AREQ US 4936679 A UPAB: 19930922

The optical fibre transducer system has an energy generator for transmitting pulsing energy at various frequencies to bidirectional couplers for each frequency. The couplers record the intensity and further transmit the pulsing energy to a wavelength multiplexer/demultiplexer. The wavelength multiplexer/demultiplexer combines the supplies into a single output for an optic fibre which includes an optical delay sufficient to time separate the pulsing waves of energy. Reflected energy is transmitted back through the same wavelength multiplexer/demultiplexer, bidirectional coupler so that the recorder intensity of transmission and reflectance are comparable with system influence.

A method is also shown for use of an optical fibre system including the components set forth and the system requires the generation and combination of the various frequencies of energy in a multiplexer/demultiplexer, the delay for time separation and the detection in a bidirectional coupler of transmitted and reflected energy.

USE - Catheter instrument.

L110 ANSWER 29 OF 29 WPIN (C) 1992 THOMPSON PERWENT

AN 1981-04/000 [16] WPIN

TI Investigating unknown substance - by comparing spectral peak table for substance with library of chemical structural units.

DC 004 S03

IN CARTER, H V; COATES, J B; FOFI, M A; HANNAH, R W; SAVITERY, A

FA (REPE) FERPIN-ELMER CORP

CYC 1

E1 GE 2070275 A 19811103 1981360* 17p

DE 4104178 A 19811116 (198149)

US 4365307 A 19821221 (198302)

GB 2076235 B 19840201 (198405)

ADT GE 2070275 A GE 1981-3088 19810202

PRAI US 1980-112337 19800207

IC G01N021-31; G06F015-20

AB GE 2076235 A UPAB: 19830915

Appts. for determining the nature of an unknown substance can enter a spectral peak table for the substance into computing appts. and adjust it to a first preselected standardised format and compare it with a first library of chemical structural units in a memory. The possible units most closely corresponding to the unknown substance are listed.

The peak table is readjusted to a second preselected standardised format. A file is formed for the unknown substance including its readjusted peak table and data of the list of possible chemical structural units is compared with files in a second library each contg. data for a

known substance and including its readjusted peak table and data of its chemical structural unit. A list of known substances most closely corresponding to the unknown substance is presented.

FS CFI EPI

FA AB

MC CFI: J04-B01A

EPI: S03-E04A9; S03-E09X

ABEQ GB 2070135 B CPAB: 19930915

Appts. for determining the nature of an unknown substance can enter a spectral peak table for the substance into computing appts. and adjust it to a first preselected standardised format and compare it with a first library of chemical structural units in a memory. The possible units most closely corresponding to the unknown substance are listed.

The peak table is readjusted to a second preselected standardised format. A file is formed for the unknown substance including its readjusted peak table and data of the list of possible chemical structural units is compared with files in a second library each contg. data for a known substance and including its readjusted peak table and data of its chemical structural unit. A list of known substances most closely corresponding to the unknown substance is presented.

= d his

(FILE 'HOME' ENTERED AT 06:39:26 ON 08 JUL 2002)

SET COST OFF

FILE 'HOMPLUS' ENTERED AT 06:39:41 ON 08 JUL 2002

E WILES T /AU

L1 1 S E6

E TURNER D/AU

L2 100 S E3,E15

L3 20 S E28,E44

L4 14 S E60,E01

L5 1 S E41

E O CORNELL M/AU

L6 1 S E6

E O CORNELL M/AU

L7 10 S E7,E4

L8 40 S E91,E18,E01

E PARMIGIANI G/AU

L9 10 S E5

E CLYDE M/AU

L10 1 S E4,E0,E7

L11 900 S (RECTON?(LO DICKIN?)/PA,CS

E MATHEMATIC/CT

E E1+ALL

L12 9040 S E1

E E1+ALL

L13 40 S E3,E5

L14 5000 S E2

L15 81203 S E1+NT

L16 171388 S MATH?

L17 12037 S L16 AND L12-L15

L18 230020 S L12-L17

E TURBIDITY/CT

E E5+ALL

L19 1000 S E1

L20 38976 S TURBID?

L21 2602 S E6+NT OR E9+NT

L22 6.6 S E1+NT

L23 1800 S E4,E1

E E9+ALL

L24 4452 S E5
 E REDOX/CT
 E E2+ALL
 L25 790 S E2, E3
 L26 5182 S E2
 L27 18308 S E2+NT
 E E2+ALL
 L28 31824 S E2+NT
 L29 187396 S E28+NT OR E70+NT OR E71+NT OR E72+NT
 L30 106788 S REDOX
 L31 540 S L18 AND L19-L24
 L32 5241 S L18 AND L25-L30
 L33 0 S L31 AND L32
 L34 5775 S L31, L32
 L35 66 S L34 AND (WAVELENGTH OR WAVE LENG?)
 E LIGHT/CT
 E E2+ALL
 L36 794 S L18 AND E2+NT
 L37 9003 S L18 AND (E23+NT OR E23+NT OR E25+NT OR E26+NT OR E27+NT OR E2
 L38 79 S L18 AND (E40+NT OR E47+NT OR E48+NT OR E49+NT OR E50+NT OR E5
 L39 6136 S L18 AND LIGHT?
 L40 2364 S L18 AND (WAVELENGTH OR WAVE LENG?)
 L41 26 S L18 AND WL
 L42 303 S L30-L41 AND L31
 L43 251 S L30-L41 AND L32
 L44 4 S L42 AND L43
 E SCREENING/CT
 E E2+ALL
 E DRUG SCREENING/CT
 E E2+ALL
 L45 18488 S E2, E2+NT
 L46 4585 S E2+NT
 L47 1 S L42, L43 AND L45, L46
 E APPARATUS/CT
 L48 0 S L43, L43 AND E3
 L49 1 S L43, L43 AND E3/CW
 E MEASURING APPARATUS/CT
 E E2+ALL
 L50 4 S L18 AND E4, E5
 L51 3112 S L18 AND E3+NT
 E E2+ALL
 L52 41 S L18 AND E3, E2+NT
 E E2+ALL
 L53 83 S L18 AND E2+NT
 L54 83 S L18 AND E3+NT
 L55 187 S L18 AND E2+NT
 L56 3875 S L42-L55
 L57 36 S L56 AND L31
 L58 37 S L56 AND L32
 L59 52 S L57, L58
 L60 1082 S L18 AND GRIN/IC, ICM, ICS
 L61 49 S L60 AND L34
 L62 33 S L61 AND L35-L59
 L63 34 S L61-L62
 L64 13 S L63 AND (3 OR 10 OR 4)/SC, SX
 L65 23 S L58 NOT L63, L64
 SEL ON AN 3 6
 L66 1 S E1-E6
 L67 49 S L63, L64, L66
 L68 31 S L1-L11 AND L13
 L69 8 S L63 AND L12-L67
 E O'CONNELL M/AU
 L70 60 S E3, E4

L71 19 S E51,E52,E60
 L72 2 S L18 AND L70,L71
 L73 8 S L68,L72 AND L19-L67
 SEL DN AN 5-8
 L74 4 S L73 NOT E1-E12
 L75 38 S L67 AND (PDC=20000531 OR PDC=20000531 OR AD<=20000531)
 L76 41 S L74,L75
 L77 41 S L76 AND L1-L76
 SEL DN AN L77 4 6 10 11 16 17 21 24-28 33 34 38 39 41
 L78 24 S L77 NOT E13-E63
 L79 24 S L78 AND (GROW? OR CONCENTR? OR LIGHT? OR WAVELENG? OR WAVE LE

FILE 'HCAPLUS' ENTERED AT 07:32:51 ON 08 JUL 2002

L80 18 S L18 AND G01H001-31/IC,ICM,ICS
 L81 14 S L80 NOT L79
 L82 10 S L81 AND (PDC=20000531 OR PDC=20000531 OR AD<=20000531)
 L83 10 S L82 AND L1-L82
 SEL L83 DN AN 2 4 5 6 12 13 14 16 17 20
 L84 10 S L83 NOT E64-E93
 L85 10 S L84 AND (GROW? OR CONCENTR? OR LIGHT? OR WAVELENG? OR WAVE LE

FILE 'WPIX' ENTERED AT 07:41:24 ON 08 JUL 2002

E WILES T/AU
 L86 1 S E4
 E O CONNELL M/AU
 L87 10 S E3,E4
 E O CONNELL M/AU
 L88 1 S E3
 E O CONNELL M/AU
 L89 9 S E3,E4
 E FARMIGIANI S/AU
 L90 2 S E3
 E TURNER D/AU
 L91 95 S E3,E13,E14
 E CLYDE M/AU
 L92 1 S E3
 E BECT/PA
 L93 2073 S (BECT?(1)DICK?) /PA
 E BECT/PACO
 E E3-ALL
 L94 2070 S E1
 L95 1380 S G01H001-31/IC,ICM,ICS
 L96 6 S L95 AND G12M001-34/IC,ICM,ICS
 L97 323 S L96 AND G01H003/IC,ICM,ICS
 L98 1 S G06F/IC,ICM,ICS AND L97
 L99 4 S L98-L94 AND L95
 L100 9 S L98,L99 AND L96,L97
 L101 9 S L100 AND G01H/IC,ICM,ICS
 SEL DN AN 4-8
 L102 4 S L101 NOT E1-E13
 L103 7 S L96,L99-L100 NOT L101
 SEL DN AN 1
 L104 1 S L103 AND E14-E15
 L105 5 S L102,L104
 L106 6 S L99,L105
 L107 25 S L95 AND G06F/IC,ICM,ICS
 L108 12 S L107 NOT L98-L106
 L109 23 S L106,L108 AND L86-L108
 L110 20 S S01/DC AND L109

FILE 'WPIX' ENTERED AT 08:01:46 ON 08 JUL 2002